Pokeweed antiviral protein alters splicing of HIV-1 RNAs, resulting in reduced virus production

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ABSTRACT
Processing of HIV-1 transcripts results in three populations in the cytoplasm of infected cells: full-length RNA, singly spliced, and multiply spliced RNAs. Rev, regulator of virion expression, is an essential regulatory protein of HIV-1 required for transporting unspliced and singly spliced viral transcripts from the nucleus to the cytoplasm. Export allows these RNAs to be packaged into virus particles. In our study, we investigate the activity of pokeweed antiviral protein (PAP), a glycosidase isolated from the pokeweed plant Phytolacca americana, on the processing of viral RNAs. We show that coexpression of PAP with a proviral clone alters the splicing ratio of HIV-1 RNAs. Specifically, PAP causes the accumulation of multiply spliced 2-kb RNAs at the expense of full-length 9-kb and singly spliced 4-kb RNAs. The change in splicing ratio is due to a decrease in activity of Rev. We show that PAP depurinates the rev open reading frame and that this damage to the viral RNA inhibits its translation. By decreasing Rev expression, PAP indirectly reduces the availability of full-length 9-kb RNA for packaging and translation of the encoded structural proteins required for synthesis of viral particles. The decline we observe in virus protein expression is not due to cellular toxicity as PAP did not diminish translation rate. Our results describing the reduced activity of a regulatory protein of HIV-1, with resulting change in virus mRNA ratios, provides new insight into the antiviral mechanism of PAP.

Keywords: HIV-1; Rev; pokeweed antiviral protein; glycosidase; ribosome inactivating protein

INTRODUCTION
The current treatment strategy for HIV-1 includes combined viral inhibitors, referred to as highly active antiretroviral therapy (HAART) (Broder 2010; Doyle and Geretti 2012). A significant challenge for treatment is the mutagenic nature of the virus, which has often led to drug resistance (Kuritzkes 2011; Tang and Shafer 2012). The administration of drug mixtures aims to reduce the chance of resistance by targeting several steps in the virus lifecycle, namely entry, reverse transcription, integration, and maturation (Flexner 2007; Mehellou and De Clercq 2010). Though HAART has significantly extended the lifespan and delayed the onset of AIDS for HIV-1-infected individuals, concerns with this treatment include side effects, resistance, and high cost. Discovery and development of additional ways to inhibit HIV-1 are a constant and essential research effort.

Pokeweed antiviral protein (PAP), a glycosidase isolated from the pokeweed plant (Phytolacca americana), removes purines from various RNA templates (Endo et al. 1988; Karran and Hudak 2008; Mansouri et al. 2009). The enzyme is of research interest because of its broad-range activity against viruses of agricultural and medical relevance. For example, the antiviral activity of PAP against HIV-1 has been observed in mammalian cell cultures and animal models. Addition of picomolar concentrations of PAP to HIV-infected CD4+ T cells substantially inhibited the synthesis of viral proteins without affecting viability or function of the cells (Zarling et al. 1990). Lack of cytotoxicity has been reported in cultured human cells (HEK 293T) (Chan Tung et al. 2008; Mansouri et al. 2009), monkeys (Macaca cynomolgus) (Uckun et al. 1998), and in mouse models (Hu-PBL-SCID, B6C3F1, and CD-1) (Uckun et al. 1998; D’Cruz et al. 2004) at concentrations of PAP that inhibit viral replication. Co-incubation of HIV-1 genomic RNA with PAP resulted in concentration-dependent release of adenines into the media, indicating depurination of the RNA (Rajamohan et al. 1999). These studies have focused primarily on evaluating the clinical potential of PAP as an anti-HIV-1 agent and have established that PAP is a potent, apparently nontoxic, inhibitor of the virus (Uckun et al. 1999).
The goal of this study was to investigate the antiviral mechanism of PAP, specifically to determine what post-transcriptional events in the HIV-1 lifecycle are affected by PAP. Following integration of the HIV-1 provirus and transcription of genomic RNA by cellular RNA polymerase, the full-length viral RNA (∼9 kb) is spliced into transcripts of ∼4 and 2 kb (Purcell and Martin 1993). The unspliced 9-kb transcripts, as well as the singly spliced 4-kb transcripts, contain introns and cis-acting repressive sequences (CRS) that promote retention in the nucleus (Schwartz et al. 1992; Zhang et al. 1996). As a result, only the 2-kb multiply spliced mRNAs exit from the nucleus using common cellular transport pathways (Kang and Cullen 1999). The 2-kb mRNAs encode three proteins, including regulator of virion expression (Rev) (Kim et al. 1989; Pomerantz et al. 1990). Rev facilitates the export of CRS-containing RNAs into the cytoplasm by binding the Rev response element (RRE) found within these transcripts (Sodroski et al. 1986; Felber et al. 1989; Cullen 1998; Pollard and Malim 1998). Although the direct role of Rev is the export of 9- and 4-kb RNAs to the cytoplasm, by virtue of its activity, Rev affects HIV-1 splicing, translation, and packaging (Hope 1999). In our current work, we show that expression of PAP alters the splicing ratio of HIV-1 RNAs by significantly reducing Rev activity. Without the important regulatory function of this protein, virus production declines.

RESULTS

Expression of PAP decreases HIV-1 protein expression without altering cellular translation rate

To characterize the anti-HIV-1 effects of PAP, 293T cells were cotransfected with the noninfectious clone pMenv(-) and the pcPAP plasmid encoding 3X-FLAG-tagged PAP. As a negative control for the catalytic activity of PAP, cells were cotransfected with pcPAPx, an active-site mutant of PAP (PE176V) that cannot depurinate RNA (Hur et al. 1995). Immunoblotting confirmed that both PAP and PAPx were effectively expressed in this system (Fig. 1A) and that PAP cotransfection with the proviral clone substantially reduced the levels of viral proteins encoded by 9- and 4-kb mRNAs, including capsid, reverse transcriptase, and Vif (Fig. 1B). Moreover, the release of particles from PAP-expressing cells was diminished, as measured by the amount of capsid protein of particles collected from growth media (Fig. 1B). This decrease agreed with our previous p24 ELISA results illustrating a 450-fold reduction in virus production from cells expressing PAP (Mansouri et al. 2012). To test whether the antiviral effect of PAP was due to cellular toxicity, cells were metabolically radiolabeled with [35S]methionine, harvested over a 6-h time period (time zero = 40 h post-transfection), and assessed for changes in translation rate, with cycloheximide treatment serving as a positive control for translation inhibition. Neither wild-type nor mutant form of PAP inhibited the overall translation rate of the cells relative to cells transfected with empty vector pcDNA3 (Fig. 1C). Mock cells were untransfected and their translation rate was always slightly higher than transfected cells.

PAP expression increases HIV-1 2-kb mRNA accumulation

To determine the effect of PAP on the level of HIV-1 mRNAs, Northern blotting was carried out using an HIV-1-specific probe that bound a region shared by all three mRNA size classes (Fig. 2A). The normal level of HIV-1 mRNAs was observed in the presence of the empty vector control, with all three mRNA classes readily detected (9, 4, and 2 kb) (Fig. 2B). PAP expression greatly altered the splicing ratio of
RRE-free and RRE-containing mRNAs within the cytoplasm (Fig. 2C), evident by the accumulation of 2-kb mRNAs and the reduction of 9- and 4-kb mRNAs (Fig. 2B). The splicing phenotype of HIV-1 remained normal with the expression of PAPx, indicating that the change seen in the presence of PAP was likely due to its catalytic activity.

**RRE-free mRNAs accumulate on polysomes**

The splicing pattern observed in PAP-expressing cells resembled that of Rev-deficient proviral clones since RRE-containing RNAs are retained in the nucleus in the absence of Rev (Pomerantz et al. 1992). The observed alteration in the splicing phenotype in the presence of PAP may be due to lack of Rev translation, even though 2-kb transcripts accumulated. To assess this possibility, polsosome analysis was carried out to identify changes in the allocation of HIV-1 mRNAs with respect to ribosome-associated and “free” pools of RNAs (i.e., those RNAs recruited to processing bodies). Northern blot analysis probing for HIV-1 mRNAs within fractions collected from density gradients revealed that there was a significant accumulation of 2-kb mRNAs on polysomes in the presence of PAP when compared with the empty vector control (Fig. 3A,B). Surprisingly, there was no significant difference in the proportion of HIV-1 mRNAs on polysomes relative to total HIV-1 mRNA in the presence or absence of PAP (Fig. 3C). This suggests that the translational profile of

![FIGURE 2. PAP increases HIV-1 splicing and 2-kb mRNA accumulation. (A) Schematic representation of the HIV-1 genome organization, transcription, and splice products. Location of cis-repressive signals (CRS) and Rev responsive element (RRE) are indicated. The annealing site for the HIV-1-specific negative-strand riboprobe is indicated below each mRNA size class as a solid line. (B) 293T cells were cotransfected with pMenv(-) proviral clone (5 μg) and pcPAP, pcPAPx, or pcDNA3 (2.5 μg). Cells were harvested 40 h post transfection and Northern blotting was performed on total cellular RNA (15 μg). Levels of the three HIV-1 mRNA classes were visualized following hybridization with the HIV-1-specific probe. Blots were also hybridized with a 5S rRNA-specific probe as a loading control. (C) The abundance of HIV-1 mRNAs in each size class was measured by quantifying the band intensity using a PhosphorImager. The ratio of RRE-free mRNAs (2 kb) to RRE-containing mRNAs (9 and 4 kb) was plotted relative to 5S rRNA. Error bars represent means ± SE for three independent experiments.

![FIGURE 3. PAP causes RRE-free mRNAs to accumulate on polysomes. (A) 293T cells were cotransfected with pMenv(-) proviral clone (5 μg) and pcPAP or pcDNA3 (2.5 μg). Cells were harvested 40 h post-transfection, and lysates were resolved by density gradient centrifugation and collected in 12 fractions. RNA was isolated from each fraction and levels of the three HIV-1 mRNA classes were visualized by Northern blotting following hybridization with the HIV-1-specific probe. The polysome profile was visualized prior to Northern blotting by ethidium bromide staining of the agarose gel. (B) The abundance of HIV-1 mRNAs in each size class was measured by quantifying the band intensity using a PhosphorImager and the ratio of RRE-free mRNAs (2 kb) to RRE-containing mRNAs (9 and 4 kb) was plotted. (C) The abundance of HIV-1 mRNAs on polysomes was plotted as a percentage of total HIV-1 RNA. Error bars in B and C represent means ± SE for three independent experiments.](http://example.com/fig2.png)
HIV-1 was not altered by PAP and the antiviral effect was not likely due to recruitment of viral mRNAs to nontranslating pools of RNA.

**Level of rev mRNA increases in the presence of PAP**

Since the accumulated 2-kb mRNAs associated with polysomes in the presence of PAP, it was possible that more regulatory proteins (Tat, Rev, and Nef) were being produced. However, the splicing pattern observed in PAP-expressing cells was similar to Rev-deficient clones, suggesting that the level of rev mRNA may be reduced in these cells. RNase protection assay was carried out to identify which 2-kb mRNAs were accumulating in the presence of PAP. Due to overlapping sequences in the three mRNA classes of HIV-1, the ~2-kb mRNA population was isolated from low-melt agarose gels and equal amounts of RNA were analyzed from cells cotransfected with pcPAP or the empty vector. Figure 4A is a schematic representation of the three 2-kb mRNAs of HIV-1, showing the annealing site of the riboprobe and the expected sizes following the assay. The housekeeping gene glyceraldehyde 3-phosphate dehydrogenase (GAPDH) was used as a loading control due to its size similarity to the 2-kb mRNAs (~1.4 kb) and thus could be cut from the gel at the same time. The levels of all three 2-kb mRNAs, tat, rev, and nef, increased approximately twofold with PAP expression (Fig. 4B), relative to cells transfected with empty vector (Fig. 4C). Therefore, rev mRNA was present in cells in which the splicing phenotype resembled a Rev-deficient clone.

**PAP decreases Rev-dependent mRNA transport**

The presence of rev mRNA in PAP-expressing cells, taken together with an increase in HIV-1 mRNA splicing ratio, suggested that the rev mRNA was not producing active Rev protein. We attempted analysis of Rev by immunoblot but could only detect Rev when overexpressed, by transfection of cells with a plasmid encoding rev. Therefore, we tested the level of Rev activity from the proviral clone by transfecting cells with the Rev activity reporter plasmid pDM128. This construct contains the chloramphenicol acetyltransferase (CAT) gene upstream of a RRE, between splice donor and acceptor sites (Fig. 5A). The CAT mRNA would only be detected in the cytoplasm if Rev facilitated its transport from the nucleus; otherwise, the CAT open reading frame would be spliced from the mRNA. Northern blot assay using a CAT-specific riboprobe showed that PAP significantly reduced the level of Rev activity relative to the empty vector and PAPx controls (Fig. 5B). In addition, Rev activity was rescued by transfecting cells with a plasmid encoding rev (pcREV) in the presence of PAP. The lack of adequate Rev activity from the proviral clone in the presence of PAP was consistent with the observed increase in the splicing phenotype of HIV-1.

**pcRev rescues the HIV-1 splicing phenotype in the presence of PAP**

To assess whether the PAP-induced change to HIV-1 mRNA splicing was due to loss of Rev activity, similar rescue experiments were performed. PAP-expressing cells were supplemented with a plasmid encoding Rev and analyzed for restoration of the splicing phenotype. Addition of pcRev regained the splicing phenotype in a Rev-deficient proviral clone pRev(-) (Fig. 6A, second lane), showing that Rev expressed from this plasmid functioned as expected. Similarly, the rescue of the splicing phenotype was successful for both HIV-1 proviral clones, pRev(-) (Fig. 6A, third lane), and
pMenv(-) (Fig. 6B, fifth lane) in the presence of PAP. Therefore, the increase in splicing observed upon expression of PAP (Fig. 2C) was due to loss of Rev, since its addition restored the RRE-containing mRNAs while decreasing the accumulation of 2-kb mRNAs. The addition of pcRev did not detectably affect the normal splicing phenotype of pMenv(-) (Fig. 6B, eighth lane).

PAP depurinates the rev open reading frame of HIV-1

The rescue of the HIV-1 splicing ratio through the addition of pcRev, taken together with an accumulation of 2-kb mRNAs (including rev) and a substantial reduction in Rev activity, suggested that PAP may exhibit its antiviral effects on HIV-1 by targeting the rev mRNA. To determine whether PAP depurinates the viral RNA, primer extension analysis was conducted on regions of the 2-kb mRNAs. Termination of the reverse transcriptase at site A8597, and to a lesser extent G8594, was reproducibly observed with RNA isolated from PAP expressing cells (Fig. 7A) but not with the empty vector. These same nucleotides were depurinated following incubation of isolated 2-kb RNAs with PAP in vitro. The location of these potential depurination sites (Fig. 7B), numbered according to the first nucleotide of the 5′ U3 region of the proviral DNA (accession no. K03455.1), were identified by sequencing the pMenv(-) proviral DNA with the same reverse primer as used for primer extension. These termination sites were located within the rev ORF, in a region that is unique to rev and does not overlap with any other ORF of the 2-kb population.

**Depurinated rev mRNA is translated inefficiently**

Given our previous results showing that elongating ribosomes stall at depurinated nucleotides (Gandhi et al. 2008), we hypothesized that depurinated rev mRNA would not translate well. To test this directly, rev ORF was transcribed in vitro and incubated with PAP prior to translation in a cell-free system. Incorporation of radiolabeled methionine during translation of PAP-treated rev mRNA was ~25% that of untreated mRNA, indicating that PAP directly reduced the synthesis of Rev protein and depurinated mRNA was a poor template for translation (Fig. 7C). The inhibitory effect of PAP on rev mRNA translation supports our initial observation that Rev activity was reduced in PAP-expressing cells, even though rev mRNA accumulated.

**FIGURE 5.** PAP decreases the level of Rev activity. (A) Schematic representation of the pDM128 reporter construct (Hope et al. 1990) and the predicted outcomes with or without Rev activity. The annealing site for the CAT-specific negative-strand riboprobe is indicated below the reporter construct as a solid line. (B) 293T cells were cotransfected with pDM128 reporter construct (5 μg), pMenv(-) proviral clone (5 μg), and pcPAP, pcPAPx, or pcDNA3 (2.5 μg), with or without pcRev (2.5 μg). Cells were harvested 40 h post-transfection and Northern blotting was performed on total cellular RNA (15 μg). Level of the cytoplasmic CAT mRNA was visualized by hybridization with the CAT-specific probe. Blots were also hybridized with a 5S rRNA-specific probe as a loading control.

**FIGURE 6.** Addition of pcRev rescues the HIV-1 splicing phenotype in the presence of PAP. 293T cells were cotransfected with (A) pRev(-) proviral clone (5 μg) or (B) pMenv(-) proviral clone (5 μg) and pcPAP, pcPAPx, or pcDNA3 (2.5 μg), with or without pcRev (2.5 μg). Cells were harvested 40 h post-transfection and Northern blotting was performed on total cellular RNA (15 μg). Levels of the three HIV-1 mRNA classes were visualized by hybridization with the HIV-1-specific riboprobe. Blots were also hybridized with a 5S rRNA-specific riboprobe as a loading control.
FIGURE 7. PAP depurinates the rev open reading frame of HIV-1 and inhibits its translation. (A) 2-kb RNAs were isolated from 293T cells cotransfected with pMenv(-) proviral clone (5 μg) and pcPAP or pcDNA3 (2.5 μg) 40 h post-transfection; 2-kb RNAs were also isolated from 293T cells transfected only with pMenv(-) proviral clone (5 μg) and incubated with purified PAP (50, 25, 10 ng) or buffer alone (0 ng). RNAs were analyzed by primer extension using HIV-1-specific radiolabeled reverse primer complementary to region 8648–8668 nt, numbered according to the first nucleotide of the 5’ U3 region of the proviral DNA (HXB2 genome, accession no. K03455.1). No RNA indicates radiolabeled reverse primer without extension template. No pMenv(-) indicates extension from RNAs of cells without transfection of pMenv(-). cDNA products were separated through 7 M urea/8% acrylamide gel and visualized using a PhosphorImager. Manual dideoxynucleotide sequencing was performed on pMenv(-) proviral clone using the same reverse primer. A8597 and G8594, potential depurination sites in the 5’ U3 region of the proviral DNA. Open reading frames are indicated as rectangles and the two depurinated nucleotides within rev mRNA are indicated below the nucleotide scale bar. (C) Rev in vitro transcript (5 μg) was incubated with or without purified PAP (25 ng) and extracted from PAP following incubation. The treated RNA was template for in vitro translation in rabbit reticulocyte lysate containing [35S]methionine, and the levels of Rev protein synthesized were measured by trichloroacetic acid precipitation and scintillation counting. The percent incorporation values were normalized to a no-RNA control reaction with the incorporation of the −PAP sample set to 100%. Error bar ± SE for three independent experiments.

DISCUSSION

Our goal was to investigate the post-transcriptional effects of PAP on HIV-1. We transfected 293T cells with a proviral clone of HIV-1 and show that PAP expression altered the splicing ratio of cytosolic HIV-1 mRNAs. Specifically, PAP increased the amount of multiply spliced 2-kb mRNAs relative to unspliced 9-kb and singly spliced 4-kb mRNAs. This accumulation of multiply spliced mRNAs was due to a PAP-dependent decrease in Rev activity. We correlate this decreased activity with depurination of the rev ORF, which inhibited translation of the protein. Depurination of viral RNAs by PAP in mammalian cells presents an interesting model for the mechanism of antiviral activity of PAP. The creation of abasic sites in viral RNAs results in templates with intact sugar phosphate backbones that would be substrates for translation. However, our previous investigations showed that depurinated viral mRNAs stalled elongating ribosomes during translation (Gandhi et al. 2008). The current decline in translational efficiency of rev mRNA treated with PAP agrees with our previous mechanistic study. By decreasing Rev levels, PAP indirectly inhibited the release of full-length genomic RNA into the cytoplasm for packaging and the synthesis of encoded proteins required for particle production.

In its role of facilitating the transport of 9- and 4-kb HIV-1 RNAs from the nucleus to the cytoplasm, Rev is essential for balancing the products of viral gene expression. In addition, Rev promotes the translation of RRE-containing viral messages. Cotransfection of HeLa cells constitutively expressing Tat with Rev and a gag reporter construct increased the amount of Gag protein relative to cells without Rev. This enhanced translation of Gag was only observed with constructs containing a RRE, suggesting that the Rev–RRE interaction was required for increased translation (D’Agostino et al. 1992). A subsequent study using a recombinant vaccinia virus system to synthesize env mRNA directly in the cytoplasm, thereby excluding nucleo-cytoplasmic transport effects, also resulted in increased Env levels in the presence of Rev (Perales et al. 2005). Therefore, Rev regulates the processing and translation of viral RNAs, and any decline in its activity would result in substantial impact on virus production.

Given the importance of Rev as a regulatory protein, cellular factors affecting Rev have been targeted for development of new antiviral agents. For example, the eukaryotic translation initiation factor 5A (eIF-5A) is a Rev cofactor, needed for nuclear export of viral mRNAs. Drugs that block the activity of enzymes required for the hypusine modification of eIF5A have been described (Hauber et al. 2005; Hoque et al. 2009). The host factor Sam68 also promotes Rev activity by functioning synergistically with Rev to increase the association of viral RNA with the cellular translation machinery. A C-terminal deletion mutant of Sam68 altered the cytoplasmic localization of RRE-containing RNA to the nuclear periphery, preventing translation of viral RNA following nuclear export (Soros et al. 2001). Recently, the drug digoxin has been shown to decrease Rev synthesis due to its effect on host factors required for splicing of 2-kb HIV-1 RNAs (Wong et al. 2013). Targeting host cell factors instead of viral proteins may provide antiviral activity against all strains of HIV-1; however, disruption of host factors essential to the cell may also have the potential for toxicity.
As with any anti-HIV-1 drug, there are questions about side effects and this is also true for PAP, a ribosome inactivating protein that is well known to depurinate ribosomal RNA (Dallal and Irvin 1978) in addition to viral targets. PAP expression in 293T cells results in 17% depurination of rRNA (Chan Tung et al. 2008), and consistent with previous results, metabolic labeling of cells showed that this level of depurination is not sufficient to decrease the cellular translation rate (Mansouri et al. 2009). We appreciate that our current study was conducted in a cell line and our future work needs to be confirmed in primary cells over a longer time frame; however, we and others have shown previously that expression or application of PAP is not toxic to cells more relevant to HIV-1 infection, such as MT-2, a lymphoblastoid T cell line, and CD4+ T cells (Zarling et al. 1990; Mansouri et al. 2012). Therefore, we suggest that changes in HIV-1 translation would not be due to a ribotoxic effect of PAP, but rather due to a more specific targeting of viral translation caused by damage to viral RNA. Translation inhibition can be achieved through recruitment of RNAs to nontranslating RNPs, such as processing bodies, which are distinct cytoplasmic foci involved in storage and/or degradation of targeted RNAs (Parker and Sheth 2007). This was not the case for the accumulated 2-kb mRNAs in the presence of PAP; there was no change in the allocation of HIV-1 mRNAs along the profile, with the accumulated mRNAs still associating with polysomes. Therefore, it is not likely that the apparent Rev deficiency was due to the recruitment of 2-kb mRNAs away from ribosomes. Moreover, there did not seem to be accumulation of RNAs in fractions associated with 40S ribosomal subunits or monosomes, suggesting that translation of the 2-kb mRNAs was not inhibited during small subunit scanning or initiation.

Our model for the antiviral activity of PAP is that depurination of viral RNA inhibits its translation, resulting in decreased viral protein levels. We show that PAP, expressed in 293T cells, removes an adenine and a guanine base from the rev ORF. We are currently analyzing how PAP targets viral RNAs and whether particular sequence or conformation of RNA, or presence of host factors, is required. The consistent feature of PAP’s enzyme activity is that purines, rather than pyrimidines, are hydrolyzed from RNA. The investigation of PAP as a potential antiviral agent must consider the possibility of the evolution of a resistant strain. The hypermutagenic nature of HIV-1 is a concern for treatment effectiveness and is the primary reason for multi-drug therapies such as HAART (Melanou and De Clercq 2010). The lack of proofreading capability of the HIV-1 RT results in a range of from two to 10 errors per round of replication of the viral genome in vivo (Preston et al. 1988; Ji and Loeb 1992). In addition, purine to purine (or pyrimidine to pyrimidine) mismatches occur more frequently than those of purine to pyrimidine (or vice versa), likely due to similarity in nucleotide shape (Preston et al. 1988). Other studies reported misincorporation of dAMP as the most frequent mutation produced by HIV-1 RT (Ji and Loeb 1992; Balzarini et al. 2001). This incorporation bias is favorable for maintaining and even creating additional PAP-susceptible sites by preserving purine bases. Furthermore, the misincorporation bias of the HIV-1 RT avoids purine to pyrimidine mutations required for PAP resistance. However, these scenarios do not preclude the possibility that mutations at sites not depurinated by PAP could alter mRNA shape such that certain regions would no longer bind PAP. We are testing the impact of mutation on shape of the 2-kb HIV-1 RNA and whether these mutations inhibit or change location of depurination on the mRNA. Our current results show that PAP targets the mRNA of Rev, a regulatory protein of HIV-1, and that inhibition of Rev activity is not due to cellular toxicity, which support continued investigation of PAP antiviral activity.

MATERIALS AND METHODS

Plasmid constructs, cell culture, and transfection

pMenv(-) is an envelope null mutant of HIV-1 (HXB2) (Sadaie et al. 1992) and was used as a source of proviral DNA in transfections (catalog #2089, NIH AIDS Reference and Reagent Program). The pRev(-) plasmid is a Rev-deficient clone of pMenv(-) generated by the destruction of a BamHI site within the rev ORF. The pcRev plasmid encodes the mature wild-type Rev protein cloned into pcDNA3 mammalian expression vector using KpnI and EcoRI restriction sites. The reporter construct pDM128 was used to assess Rev activity and was kindly provided by Dr. A. Cochrane and has been described previously (Hope et al. 1990). The plasmids encoding 3X-FLAG-tagged mature PAP protein, pcPAP, and the active site mutant (E176V) of PAP, pcPAPx, have been described previously (Mansouri et al. 2009). Human embryonic kidney (HEK)-293T cells were maintained in Dulbecco’s Modified Eagle’s Medium (DMEM) supplemented with 10% Fetal Bovine Serum (FBS), penicillin (100 U/mL), and streptomycin (100 μg/mL). Cells were grown on 10-cm treated plates in a humidified incubator at 37°C with 5% CO2. Plasmid DNA was transfected into cells (5 × 105 cells/mL) by standard calcium phosphate coprecipitation (Chen and Okayama 1988). Cells were refed 18 h after transfection and were harvested 40 h after transfection. Virus particles released from cells were isolated from 1 mL of medium 40 h after transfection. The medium was centrifuged at 5000g for 5 min at room temperature to remove cell debris, and the supernatant was centrifuged at 16,100g for 90 min at 4°C to pellet virus particles.

Polysome analysis

To determine the effects of PAP on the translational profile of HIV-1, polysome analysis was carried out. HEK 293T cells were transfected, and 40 h following transfection the growth medium was replaced with 10 mL DMEM containing 100 μg/mL cycloheximide and cells were incubated for 20 min at 37°C. Cells were then harvested in 1X PBS containing 100 μg/mL cycloheximide and 1 mM PMSF and centrifuged at 1000g for 5 min at 4°C, washed with 10 mL of ice cold 1X PBS containing 100 μg/mL cycloheximide and centrifuged again. The resulting pellets were resuspended in 0.5 mL Magnesium
Buffer (10 mM HEPES-KOH, 10 mM NaCl, 7 mM MgCl₂, 3 mM CaCl₂, 1 mM DTT, 0.5% NP-40), vortexed for 10 sec and kept for 10 min on ice. Samples were centrifuged at 14,000g for 4 min at 4°C, the supernatant was combined with 1 mg of heparin and quantified at OD₅₅₀nm. Equal amounts (40 OD₅₅₀nm) were loaded onto 14%–47% linear sucrose gradients and centrifuged at 240,000g for 2.5 h at 4°C. The gradients were divided into 12 fractions and incubated in 1% SDS and 0.48 mM proteinase K for 30 min at 37°C. RNA was isolated from each fraction for Northern blot analysis.

**Protein analyses**

Pellets of harvested cells were combined with equal volumes of Lysis Buffer (25 mM HEPES-KOH at pH 7.5, 2 mM EGTA, 1 mM DTT, 10% glycerol, 1% NP-40), vortexed for 20 sec, and incubated on ice for 10 min. Samples were centrifuged at 16,100g for 10 min at 4°C and total protein of the resulting supernatant was quantified by the Bradford Assay. Cell lysates or virus particles were separated through a 12% SDS–polyacrylamide gel, transferred to nitrocellulose, blocked in 5% milk and probed with appropriate primary and secondary antibodies. PAP and PAPx were detected with an anti-FLAG monoclonal antibody (1:1,000; Sigma Aldrich). HIV-1 proteins were detected with antibodies obtained through the NIH AIDS Research and Reference Reagent Program. HIV-1 Gag was detected with an anti-p24 monoclonal antibody (1:1,000; Sigma Aldrich). HIV-1 proteins were detected with antibodies obtained through the NIH AIDS Research and Reference Reagent Program. HIV-1 Gag was detected with an anti-p24 monoclonal antibody (1:1,000; Sigma Aldrich). HIV-1 RT was detected with an anti-p51 polyclonal antibody (1:5,000; NIH, cat # 3537); RT was detected with an anti-p31 polyclonal antibody (1:2,500; NIH, cat # 6195); and Vif was detected with an anti-Vif monoclonal antibody (1:2,500; Sigma Aldrich). Secondary antibodies were conjugated to horseradish peroxidase and proteins were detected by chemiluminescence.

To assess the rate of cellular translation during PAP expression, cells were metabolically radiolabeled by [35S]methionine incorporation. The day after transfection, cells were seeded in 6-well plates at a density of 1 × 10⁶ and incubated overnight at 37°C with 5% CO₂. On the following day, growth medium was replaced with 2 mL of minimum essential medium (MEM) supplemented with 10% FBS and incubated for 30 min at 37°C. Following incubation, the medium was replaced with 1 mL MEM containing 12.5 μCi of [35S]methionine (Perkin Elmer). At this time (40 h post-transfection), the negative control sample was also supplemented with 25 μg/mL cycloheximide. Cells were harvested at the indicated time points (30, 60, 120, 240, and 360 min). RIPA Buffer (60 μL; 150 mM NaCl, 10 mM HEPES-KOH at pH 7.4, 1% Triton-X100, 0.1% SDS), supplemented with 1X complete protease inhibitor cocktail tablets (Roche), was added to cell pellets, vortexed for 20 sec, and incubated on ice for 10 min. Proteins were precipitated in trichloroacetic acid and collected on glass microfiber filters. The incorporated radioactivity was quantified using a scintillation counter.

To determine the direct effect of PAP on viral RNA translation, _rev_ transcript was incubated with purified PAP and translated in a cell-free system. The _rev_ open reading frame was subcloned into pET28a at NcoI and EcoRI sites and the resulting pET-Rev plasmid was linearized with EcoRI. Run-off in vitro transcripts (5 μg) were incubated with or without 25 ng purified PAP in 1X ribosome inactivating protein buffer (60 mM KCl, 10 mM Tris-HCl at pH 7.4, 10 mM MgCl₂) for 30 min at 30°C. Reactions were extracted with phenol/chloroform/isoamyl alcohol and ethanol precipitated. The recovered transcripts were supplied as template for in vitro translation with rabbit reticulocyte lysate (Promega) in the presence of [35S]methionine for 90 min at 30°C. Protein product was precipitated with trichloroacetic acid and filtered through glass fiber filters. [35S] Incorporation was measured by scintillation counting.

**RNA analyses**

Harvested cell pellets were lysed with an equal volume of Lysis Buffer (25 mM HEPES-KOH at pH 7.5, 2 mM EGTA, 1 mM DTT, 10% glycerol, 1% NP-40) and centrifuged at 16,100g for 10 min at 4°C to pellet cellular debris. Total RNA was extracted with TRI Reagent (Molecular Research Center) following the manufacturer’s instructions and precipitated in ethanol. Northern blot analysis was used to detect and quantify levels of HIV-1 mRNAs. Isolated RNA samples (15 μg) were denatured at 85°C in formamide buffer (60% formamide, 20% formaldehyde, 20 mM MOPS at pH 7.0, 2 mM NaOAc, 1 mM EDTA) and separated through a 0.8% agarose denaturing gel. Following separation, RNA was transferred to a positively charged nylon membrane and probed for HIV-1 mRNAs with 1 × 10⁶ cpm [α-33P]-radiolabeled riboprobe specific to nucleotides 8881–9060 of pMenv(-) proviral clone. To detect chloramphenicol acetyl-transferase (CAT) mRNA for Rev activity assay, the blot was probed with a radiolabeled riboprobe specific to nucleotides 220–420 of CAT mRNA. Antisense riboprobe specific for nucleotides 24–96 of S5 RNA was used to probe the rRNA as a loading control in Northern blotting. Radiolabeled RNA was visualized and quantified with a PhosphorImager.

To isolate 2-kb mRNAs for RNase protection assay and primer extension, total RNA was separated through a 1% low-melt agarose gel. The RNA was stained with ethidium bromide and regions corresponding to 2 kb in size were cut from the gel. Gel pieces were heated for 5 min at 70°C, combined with an equal volume of 2X PK Buffer (300 mM NaCl, 25 mM EDTA, 20 mM Tris–HCl at pH 8.0, 2% SDS) and proteinase K to a final concentration of 0.25 mg/mL, and incubated for 10 min at 37°C. The RNA was extracted from the melted gel with phenol (saturated with 50 mM NaOAc at pH 5.2, 10 mM EDTA), further extracted with phenol/chloroform: isoamyl alcohol (25:24:1), and precipitated in ethanol.

The levels of 2-kb HIV-1 mRNAs were measured by RNase protection assay using an antisense riboprobe specific to nucleotides 5845–6044 of the pMenv(-) proviral clone. GAPDH mRNA was used as a loading control and was detected with an antisense riboprobe specific for nucleotides 1–341 of pTRI-GAPDH plasmid. Isolated RNA samples (15 μg) were combined with 21 μL Hybridization Buffer (400 mM NaCl, 40 mM PIPES at pH 6.4, 1 mM EDTA, 80% formamide) containing 5 × 10⁶ cpm [α-33P]-radiolabeled probe. Samples were incubated for 30 min at 80°C and then hybridized overnight at 50°C. Each sample was treated with 144 units of RNase T1 and 0.72 units of RNase A in 200 μL of RNAse Buffer (300 mM NaCl, 10 mM EDTA, 10 mM Tris–HCl at pH 7.5) for 30 min at room temperature. Samples were then incubated with 17 μL of proteinase K (10% SDS) for 30 min at 37°C. The RNA was extracted with TRI Reagent (Molecular Research Center) and chloroform and precipitated in ethanol. RNA pellets were resuspended in formamide loading buffer and separated through a 7-M urea/6% polyacrylamide gel. An RNA ladder was visualized by ethidium bromide staining and the sizes of protected radiolabeled RNA probe were detected and quantified with a PhosphorImager.
Primer extension was used to detect depurination sites of rev mRNA by PAP. The HXB28648 (5′-CTTATTGTTCTGACTCCAA TACTGTAGGAC-3′) reverse primer, used to show a depurination site in rev ORF, annealed 51 nucleotides downstream from nucleotide A8597. Isolated 2-kb HIV-1 RNAs (5 pg) were combined with 5 × 10^5 cpm of [γ-32P]-end-labeled primer, heat denatured at 90°C for 5 min, then chilled on ice. Samples were brought to 20 μL in 1× buffer (75 mM KCl, 50 mM Tris-HCl at pH 8.3, 10 mM DTT, 3 mM MgCl2, 1 mM dNTPs, 40 units of RNase inhibitor) and incubated for 10 min at room temperature. After annealing, samples were heat-denatured to 48°C and 100 units of MMLV Reverse Transcriptase enzyme (New England Biolabs) were added to each sample and incubated for 5 min, then chilled on ice. Samples were brought to 20 μL in 1× buffer and incubated at 75°C for 15 min. The PCR products were separated on a 7-M urea/8% polyacrylamide gel. To identify the depurinated nucleotide, dideoxynucleotide sequencing of pMenv(-) HIV-1 plasmid was performed with the same reverse primer used for the primer extension assay. Radiolabeled cDNAs were visualized with a PhosphorImager.

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