Tip110 Protein Binds to Unphosphorylated RNA Polymerase II and Promotes Its Phosphorylation and HIV-1 Long Terminal Repeat Transcription*

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Background: Tip110 synergizes with HIV-1 Tat protein and transactivates Tat-mediated HIV-1 LTR promoter, but the underlying mechanisms have not been understood.

Results: Tip110 bound to unphosphorylated RNApolII and led to increased P-TEFb recruitment and RNApolII phosphorylation, enhancing Tat-mediated transcription elongation of the HIV-1 LTR.

Conclusion: Tip110 is involved in transcription regulation of the HIV-1 LTR promoter.

Significance: These findings provide additional insights into HIV-1 transcription.

Transcription plays an important role in both HIV-1 gene expression and replication and mandates complicated but coordinate interactions between the host and virus. Our previous studies have shown that an HIV-1 Tat-interacting protein of 110 kDa, Tip110, binds to and enhances Tat function in Tat-mediated HIV-1 gene transcription and replication (Liu, Y., Li, J., Kim, B. O., Pace, B. S., and He, J. J. (2002) J. Biol. Chem. 277, 23854–23863). However, the underlying molecular mechanisms by which this takes place were not understood. In this study, we demonstrated that Tip110 bound to unphosphorylated RNA polymerase II (RNApolII) in a direct and specific manner. In addition, we detected Tip110 at the HIV-1 long terminal repeat (LTR) promoter and found that Tip110 expression was associated with increased phosphorylation of serine 2 of the heptapeptide repeats within the RNApolII C-terminal domain and increased recruitment of positive transcription elongation factor b to the LTR promoter. Consistent with these findings, we showed that Tip110 interaction with Tat directly enhanced transcription elongation of the LTR promoter. Taken together, these findings have provided additional and mechanistic evidence to support Tip110 function in HIV-1 transcription.
lated protein, Tat-SF1, which interacts with large RNAPII elongation-splicing complexes as well as the Tat-P-TEFb complex and stimulates HIV-1 transcription (22–24).

Tat-interacting protein of 110 kDa (Tip110), also known as SART3/p110, was initially identified as a nuclear protein in the mRNA splicing process during the study of human U6 snRNA capping enzyme (25). It was later reported to be a human homologue of yeast protein Prp24 and associates only transiently with U6 and U4/U6 snRNP during the recycling phase of the spliceosome cycle (26, 27). Human Tip110 was found to encode a tumor rejection antigen in various cancers; it is expressed in a number of cancer cell lines as well as the majority of cancer tissues (28–30). In addition, Tip110 also binds to the androgen receptor through the nuclear receptor box and functions as a repressor of androgen receptor transcription activation through interference with the complex formation between the androgen receptor and androgen receptor-responsive element (31).

We have previously reported that Tip110 is important for HIV-1 gene expression and virus replication by interacting with Tat and enhancing Tat-mediated transactivation activity (32). However, the molecular mechanisms underlying this process were not understood. In the current study, we characterized Tip110 interaction with RNAPII and P-TEFb and its effects on RNAPII phosphorylation and HIV-1 transcription initiation and elongation. We showed that Tip110 bound to unphosphorylated RNAPII and increased P-TEFb recruitment to the TAR-Tat-P-TEFb transcription complex, which was associated with increased phosphorylation of serine 2 of the heptapeptide repeats within the RNAPII CTD and enhanced Tat-mediated transcription elongation of the LTR promoter.

**MATERIALS AND METHODS**

**Cell Lines and Cell Transfections—**293T cells were purchased from the American Tissue Culture Collection, and U373MAGI and CEM-GFP cells were obtained from the National Institutes of Health AIDS Research and Reference Reagent Program. The cells were maintained in either Dulbecco’s modified Eagle’s medium (293T and U373MAGI), or RPMI 1640 medium (CEM-GFP), with 10% fetal bovine serum and incubated at 37 °C in 10% CO₂. Transfections were carried out using the standard calcium phosphate precipitation method. pcDNA3 was used to equalize the total amount of DNA, whereas pEGFP was used to ensure that transfection efficiencies were comparable among all transfections.

**Plasmids—**Construction of pTip110.His, pGEX-Tip110, pTip110.HA, Tip110 deletion mutants (ΔRRM, ΔNLS, and ΔCT), pTat.Myc, and pLTR-Luc plasmid was described previously (32). GST-CTD plasmid was a gift from Dr. David Price (University of Iowa, Iowa City, IA). Escherichia coli (E. coli) BL21 cells. The culture was grown to an A₆₀₀ of 0.6. At this point, 0.5 mM isopropyl-β-thiogalactoside was added, and the culture was incubated at 37 °C for an additional 3 h. The bacteria were first lysed with a lysis buffer (50 mM Tris-HCl, pH 7.9, 12.5 mM MgCl₂, 0.5 mM EDTA, 100 mM KCl, 20% glycerol, 1 mM β-mercaptoethanol, 10 μM ZnCl₂) and then disrupted in a French press. The supernatant was collected and incubated with glutathione-Sepharose 4B, and the protein was eluted with 4 mM reduced glutathione and recovered by centrifugation and then washed with PBS five times, and the bound proteins were eluted by adding glutathione beads to remove the GST protein. For the GST pull-down assay, GST-CTD (6 μg) protein was first phosphorylated overnight at 30 °C with 6 μl of casein kinase I (New England Biolabs) and 1 mM ATP with the phosphorylation buffer provided with the enzyme (New England Biolabs). Meanwhile, 4 μg of GST-Tip110 protein was digested with 1 μl of thrombin at 30 °C overnight, and then Tip110 was purified by adding glutathione beads to remove the GST protein. For the GST pull-down assay, GST-CTD (6 μg) protein was first phosphorylated overnight at 30 °C with 6 μl of casein kinase I (New England Biolabs) and 1 mM ATP with the phosphorylation buffer provided with the enzyme (New England Biolabs). Meanwhile, 4 μg of GST-Tip110 protein was digested with 1 μl of thrombin at 30 °C overnight to remove the GST tag. Unphosphorylated or phosphorylated GST-CTD proteins were then immobilized onto 30 μl of glutathione beads at 4 °C for 2 h. Then the protein-bound beads were incubated with purified Tip110 protein in 500 μl of GST pull-down buffer (20 mM HEPES, pH 7.9, 150 mM NaCl, 0.5 mM EDTA, 10% glycerol, 0.1% Triton X-100, 1 mM DTT) at room temperature for 2 h. Subsequently, the protein-bound beads were washed with PBS five times, and the bound proteins were eluted from the beads by 4× SDS-PAGE sample buffer. The proteins were separated by SDS-PAGE and analyzed by immunoblotting.

**ChIP Assay—**Cells (5 × 10⁶) were first cross-linked with 1% formaldehyde at room temperature for 20 min; the cross-linking was terminated by the addition of glycine to a final concentration of 0.125 M. The cells were washed by PBS three times, and the cell pellet was suspended in a cell lysis buffer (85 mM KCl, 0.5% Nonidet P-40, 5 mM HEPES, pH 8.0) and then incubated on ice for 10 min. The nuclei were recovered by centrifugation and centrifuged 72 h post-transfection in a cell lysis buffer (50 mM Tris-HCl, pH 8.0, 0.5% Nonidet P-40, 2 mM EDTA, 137 mM NaCl, 10% glycerol, 0.5% sodium deoxycholate, 0.2% sodium azide, 0.004% sodium fluoride, 1X protease inhibitor mixture, 1 mM sodium orthovanadate, pH 7.25). Whole cell extracts were obtained by centrifugation at 15,000 × g for 10 min. Whole cell extracts of an equal amount of protein were separated by 8–12% SDS-PAGE and then electrotransferred to HyBond-P membrane (Amersham Biosciences). The membrane was probed with primary antibodies and appropriate peroxidase-labeled secondary antibody and then visualized with a homemade ECL system. For immunoprecipitation, whole cell extracts of 500 μg of protein were first pre-cleared using 20 μl of protein A-agarose beads (Millipore) and then incubated by rotation with 1 μg of antibody and 60 μl of protein A-agarose beads at 4 °C overnight. The beads were recovered by centrifugation and then washed with a washing buffer (50 mM Tris-HCl, pH 8.0, 0.5% Nonidet P-40, 2 mM EDTA, 0.4 mM NaCl, 10% glycerol) four times. The beads were suspended in a SDS-PAGE sample buffer and used for SDS-PAGE and Western blot analysis.

**Whole Cell Lysate Preparation, Immunoprecipitation, and Western Blot Analysis—**Unless stated otherwise, cells were harvested 72 h post-transfection in a cell lysis buffer (50 mM Tris-HCl, pH 8.0, 0.5% Nonidet P-40, 2 mM EDTA, 137 mM NaCl, 10% glycerol, 0.5% sodium deoxycholate, 0.2% sodium azide, 0.004% sodium fluoride, 1X protease inhibitor mixture, 1 mM sodium orthovanadate, pH 7.25). After incubation on ice for 20 min, whole cell extracts were obtained by centrifugation at 15,000 × g for 10 min. Whole cell extracts of an equal amount of protein were separated by 8–12% SDS-PAGE and then electrotransferred to HyBond-P membrane (Amersham Biosciences). The membrane was probed with primary antibodies and appropriate peroxidase-labeled secondary antibody and then visualized with a homemade ECL system. For immunoprecipitation, whole cell extracts of 500 μg of protein were first pre-cleared using 20 μl of protein A-agarose beads (Millipore) and then incubated by rotation with 1 μg of antibody and 60 μl of protein A-agarose beads at 4 °C overnight. The beads were recovered by centrifugation and then washed with a washing buffer (50 mM Tris-HCl, pH 8.0, 0.5% Nonidet P-40, 2 mM EDTA, 0.4 mM NaCl, 10% glycerol) four times. The beads were suspended in a SDS-PAGE sample buffer and used for SDS-PAGE and Western blot analysis.

**Recombinant Protein Purification and GST Pull-down Assay—**pGEX-Tip110 and pGST-CTD were first transformed into Escherichia coli BL21 cells. The culture was grown to an A₆₀₀ of 0.6. At this point, 0.5 mM isopropyl-β-thiogalactoside was added, and the culture was incubated at 37 °C for an additional 3 h. The bacteria were first lysed with a lysis buffer (50 mM Tris-HCl, pH 7.9, 12.5 mM MgCl₂, 0.5 mM EDTA, 100 mM KCl, 20% glycerol, 1 mM β-mercaptoethanol, 10 μM ZnCl₂) and then disrupted in a French press. The supernatant was collected and incubated with glutathione-Sepharose 4B, and the protein was eluted and concentrated. GST-Tip110 protein was digested with thrombin at 30 °C overnight, and then Tip110 was purified by adding glutathione beads to remove the GST protein. For the GST pull-down assay, GST-CTD (6 μg) protein was first phosphorylated overnight at 30 °C with 6 μl of casein kinase I (New England Biolabs) and 1 mM ATP with the phosphorylation buffer provided with the enzyme (New England Biolabs). Meanwhile, 4 μg of GST-Tip110 protein was digested with 1 μl of thrombin at 30 °C overnight to remove the GST tag. Unphosphorylated or phosphorylated GST-CTD proteins were then immobilized onto 30 μl of glutathione beads at 4 °C for 2 h. Then the protein-bound beads were incubated with purified Tip110 protein in 500 μl of GST pull-down buffer (20 mM HEPES, pH 7.9, 150 mM NaCl, 0.5 mM EDTA, 10% glycerol, 0.1% Triton X-100, 1 mM DTT) at room temperature for 2 h. Subsequently, the protein-bound beads were washed with PBS five times, and the bound proteins were eluted from the beads by 4× SDS-PAGE sample buffer. The proteins were separated by SDS-PAGE and analyzed by immunoblotting.
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FIGURE 1. No complex formation between Tip110 and P-TEFb. A, 293T cells were transfected with pTip110.HA (lane 1) or pcDNA3 (lane 2). Whole cell extracts (WCE) were prepared and immunoprecipitated (IP) with α-HA, α-cyclin T1 (CycT1), or α-CDK9 antibody, followed by Western blot analysis (WB) with α-HA, α-CycT1, or α-CDK9 antibody. B, WCE were prepared from 293T cells and immunoprecipitated with α-Tip110 antibody or an isotype-matched IgG, followed by Western blot analysis with α-CycT1, α-CDK9, or α-Tip110 antibody. C, WCE were prepared from 293T cells and immunoprecipitated with α-CycT1 antibody or an isotype-matched IgG, followed by Western blot analysis with α-Tip110 or α-CDK9 antibody.

RESULTS

Tip110 Did Not Complex with Cyclin T1 and CDK9—Our previous work has shown that Tip110 enhances HIV-1 replication by activating Tat-mediated LTR transcription (32). Because Tip110 and P-TEFb co-localize within nuclear speckle structures (35) and P-TEFb is responsible for HIV-1 transcription elongation, we first determined whether Tip110 interacted with CDK9 or cyclin T1 by immunoprecipitation and Western blot analysis. The primers used to amplify the short G-less transcripts were 5′-GGG TCT CTC AGG AGG GAG-3′ and 5′-AAA ACC AAA CCC TGC GCT CCA TCG CCA-3′. The primers used to amplify the long G-less transcripts were 5′-GGC TTA AGC AGT GGG TTC CCT AG-3′. The PCR program consisted of 35 cycles of 94 °C for 1 min, 55 °C for 1 min, and 72 °C for 30 s.

In Vitro Transcription Assay—An in vitro transcription assay was performed as described previously (33) with some minor modifications. Briefly, recombinant Tat protein and Tip110 protein were mixed in buffer D (20 mM HEPES, pH 7.9, 0.1 mM KCl, 20% glycerol, 0.2 mM EDTA, 0.5 mM DTT) and added to 3.5 µl of nuclear extracts (Promega) on ice. Then 200 ng of linearized HIV-1 dG-less DNA, unlabeled NTP, 4 units of RNase inhibitor, and 400 mM sodium citrate were added with a total 25-µl transcription reaction. The mixture was incubated at 30 °C for 30 min, followed by treatment with 1 µl of DNase I to remove the DNA template. RNA transcripts were recovered by phenol extraction and ethanol precipitation, and the levels of long G-less transcripts were quantitated by RT-PCR, followed by the densitometric analysis of the RT-PCR products; the ratio of the PCR products derived from the long G-less transcripts was 5′-GGC GAG TAC TTC AAG AAC TGC-3′ and 5′-AAA ACC AAA CCC TGC GCT CCA TCG CCA-3′. The levels of long and short transcripts were quantitated by RT-PCR, followed by the densitometric analysis of the RT-PCR products; the ratio of the PCR products derived from the short transcripts was calculated and used to express the elongation efficiency.

gation at 3,000 × g for 5 min and then suspended in a nuclear lysis buffer (10 mM EDTA, 1% SDS, 50 mM Tris-HCl, pH 8.1). The nuclei were incubated on ice for an additional 10 min, and the supernatants were collected by centrifugation at 15,000 × g for 10 min and saved as nuclear lysates. The nuclear lysates were then sonicated on ice with 10 pulses, each for 15 s, to generate chromatin DNA with an average size of 600 bp. The sonicated DNA was diluted 10-fold with a buffer (165 mM NaCl, 0.01% SDS, 1% Triton X-100, 16.7 mM Tris-HCl, pH 8.0) and pre cleared with 30 µl of protein A-Sepharose beads. The lysates were first incubated with the indicated antibodies overnight, and then 60 µl of protein A-Sepharose beads were added, and the lysates were incubated for an additional 4 h. The immunocomplexes were washed twice with a low salt buffer (150 mM NaCl, 0.1% SDS, 1% Nonidet P-40, 1 mM EDTA, 50 mM Tris-HCl), twice with a high salt buffer (500 mM NaCl, 0.1% SDS, 1% Nonidet P-40, 1 mM EDTA, 50 mM Tris-HCl), twice with a LiCl buffer (250 mM LiCl, 0.1% SDS, 1% Nonidet P-40, 1 mM EDTA, 50 mM Tris-HCl), and finally twice with TE buffer (0.25 mM EDTA, 10 mM Tris-HCl). The recovered beads were eluted with 120 µl of elution buffer (1% SDS, 100 mM NaHCO3), the supernatants were collected and incubated at 65 °C overnight to reverse the formaldehyde cross-linking. The DNA from the supernatants was recovered by phenol extraction followed by ethanol precipitation and analyzed using PCR with primers spanning the HIV-1 LTR promoter (5′-CAT CCG GAG TAC TTC AAG AAC TGC-3′ and 5′-GGC TTA AGC AGT GGG TTC CCT AG-3′) or GAPDH (5′-GAA GGTGAA GGT CGGAGT-3′ and 5′-GAA GAT GGT GAT GGG ATT TC-3′). The PCR program consisted of 35 cycles of 94 °C for 1 min, 55 °C for 1 min, and 72 °C for 30 s.

In Vitro Transcription Assay—An in vitro transcription assay was performed as described previously (33) with some minor modifications. Briefly, recombinant Tat protein and Tip110 protein were mixed in buffer D (20 mM HEPES, pH 7.9, 0.1 mM KCl, 20% glycerol, 0.2 mM EDTA, 0.5 mM DTT) and added to
endogenous Tip110 associated with P-TEFb by immunoprecipitation. We found that neither endogenous cyclin T1 nor CDK9 could be detected in the anti-Tip110 immunoprecipitates (Fig. 1B). Similarly, we did not detect Tip110 protein in either anti-CDK9 or anti-cyclin T1 immunoprecipitates (Fig. 1, C and D). Therefore, these results suggest that Tip110 did not associate with CDK9 or cyclin T1.

**Tip110 Bound to Unphosphorylated but Not Phosphorylated RNAPII**—One of the essential events in HIV-1 LTR promoter transcriptional activation is phosphorylation within the heptapeptide repeats of the RNAPII CTD at serine 2 (CTD02) and serine 5 (CTD05). To investigate whether Tip110 directly interacted with RNAPII, 293T cells were transfected with pTip110-HA (Fig. 2A). An immunoprecipitation was performed for the unphosphorylated RNAPII (CTDa), serine 2-phosphorylated RNAPII (CTD02), or serine 5-phosphorylated RNAPII (CTD05) by three highly specific antibodies (8WG16 (CTDa), H5 (CTD02), and H14 (CTD05), respectively), followed by Western blot analysis using anti-HA antibody. The results demonstrated that only CTDa, CTDo2, and CTDo5 or from different efficiencies of the antibodies for these three RNAPII forms. To ascertain Tip110 interaction with CTDa, an in vitro GST pull-down assay was performed. First, recombinant GST-CTDa protein was purified from E. coli and then phosphorylated by casein kinase I in vitro. GST-CTDa and its phosphorylation by casein kinase I (GST-CTDo) were confirmed by Western blot analysis (Fig. 3A). Meanwhile, recombinant GST-Tip110 was expressed and purified (Fig. 3B, lane 2), treated with thrombin to remove the GST tag (Fig. 3B, lane 3), and then purified using the glutathione–beads to produce recombinant Tip110 protein (Fig. 3B, lane 4). Then recombinant proteins GST-CTDa or GST-CTDo and Tip110 were used in the GST pull-down assay. The proteins that remained on the beads (bound) as well as the proteins that were present in the supernatants (unbound) were analyzed by SDS-PAGE followed by Western blot analysis using anti-Tip110 or GST antibody. The results showed direct binding of Tip110 to CTDa but not to GST protein or CTD05.

The results of these Western blots indicated that only CTDa, not CTDo2 and CTDo5, was detected to complex with Tip110 (Fig. 2). This difference could result from different expression levels of CTDa, CTDo2, and CTDo5 or from different efficiencies of the antibodies for these three RNAPII forms. To ascertain Tip110 interaction with CTDa, an in vitro GST pull-down assay was performed. First, recombinant GST-CTDa protein was purified from E. coli and then phosphorylated by casein kinase I in vitro. GST-CTDa and its phosphorylation by casein kinase I (GST-CTDo) were confirmed by Western blot analysis (Fig. 3A). Meanwhile, recombinant GST-Tip110 was expressed and purified (Fig. 3B, lane 2), treated with thrombin to remove the GST tag (Fig. 3B, lane 3), and then purified using the glutathione–beads to produce recombinant Tip110 protein (Fig. 3B, lane 4). Then recombinant proteins GST-CTDa or GST-CTDo and Tip110 were used in the GST pull-down assay. The proteins that remained on the beads (bound) as well as the proteins that were present in the supernatants (unbound) were analyzed by SDS-PAGE followed by Western blot analysis using anti-Tip110 or GST antibody. The results showed direct binding of Tip110 to CTDa but not to GST protein or CTD05.

To determine which functional domains of Tip110 interacted with CTDa, we took advantage of a series of Tip110
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FIGURE 3. Direct binding of Tip110 to RNAPII CTDa. A. GST-CTDa protein was expressed and purified from E. coli and phosphorylated by casein kinase I (CK) in vitro. GST-CTDa and CK-phosphorylated GST-CTD (GST-CTDo) were confirmed by Western blot analysis with α-CTDa, α-CTDo2, α-CTDo5, or α-GST antibody. Western blot analysis against GST was included as a loading control. B, GST-Tip110 protein was expressed and purified from E. coli (lane 2). Its GST tag was removed by thrombin digestion (lane 3), followed by glutathione bead affinity purification to remove the GST tag (lane 4). All of those proteins were subjected to Western blot analysis with α-Tip110 antibody or α-GST antibody. WCE from 293T cells were included as a control (lane 1). C, GST-CTDa, CK-phosphorylated GST-CTD (GST-CTDo), and GST proteins were immobilized with 50 μl of glutathione beads and then incubated with purified Tip110 protein. The beads and the supernatants were separated by centrifugation and collected to represent bound and unbound fractions, respectively. The beads were then subject to repetitive washes to remove unbound proteins. Then the proteins bound on the beads and in the supernatants were detected by Western blot analysis with anti-GST antibody to ensure GST binding to the glutathione beads.

WT and ΔNT were co-transfected (WT + ΔNT) and analyzed similarly. The results showed that only WT and not ΔNT was detected in the CTDa immunoprecipitates of the co-transfection (Fig. 4D). These results suggest that the N-terminal domain of Tip110 is directly involved in Tip110-CTDa interactions.

More P-TEFb Recruitment to the LTR Transcription Complex by Tip110 and Tat—Tip110 has been shown to directly interact with HIV-1 Tat protein (32). We next determined whether Tip110 interaction with Tat would recruit more P-TEFb to the HIV-1 LTR transcription complex. 293T cells were transfected with pLTR-Luc, pTat.Myc, and pTip110.HA. At 48 h post-transfection, we performed Western blot analysis to determine Tip110, Tat, CDK9, and cyclin T1 expression and found that neither Tip110 nor Tat altered the overall levels of CDK9 and cyclin T1 expression (Fig. 5A). An immunoprecipitation was performed for Tat, followed by Western blot analysis using cyclin T1 antibody. As expected, cyclin T1 was detected in anti-Myc immunoprecipitates, confirming cyclin T1/Tat interaction (Fig. 5B). Moreover, more cyclin T1 was detected in anti-Myc immunoprecipitates of cells co-expressing both Tat and Tip110 than in the cyclin T-Tat complex formed in the cells expressing Tat alone; the increase is estimated to be about 2.5-fold. Similarly, about 1.4-fold more CDK9 was detected in CTDa immunoprecipitates of cells co-expressing both Tat and Tip110 than in the CDK9-CTDa complex in the cells expressing Tat alone (Fig. 5C). To determine whether Tip110 binding to Tat/CTDa is directly involved in the increased recruitment, ΔNT that does not bind to CTDa (Fig. 4) or Tat (32) was transfected alone or with Tat; similar experiments were then performed (Fig. 5A). The results showed that unlike WT, ΔNT showed no changes in the levels of cyclin T1 and CDK9 in the transcription complex. The results were reproducible (Fig. 5D) and suggest that the interaction of Tip110 with Tat/CTDa led to the recruitment of more P-TEFb to the transcriptional complex.

Tip110 and Tat Enhanced RNAPII Phosphorylation—The most critical step during HIV-1 LTR promoter transactivation is the phosphorylation of RNAPII (36, 37). CDK7 phosphorylates serine 5 of RNAPII during the initiation of transcription, whereas CDK9 phosphorylates serine 2 during transcription elongation (4, 38). Because Tip110 could recruit more P-TEFb to

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the transcription complex, we next determined the effects of Tip110 and Tat expression on RNAPII phosphorylation. We transfected 293T cells with an increasing amount of pTip110.HA or pTat.Myc or the same amount of pTat.Myc with an increasing amount of Tip110.HA. After 48 h, Western blot analysis was performed against CTDa, CTDo2, and CTDo5. When cells were transfected with Tip110 alone, both CTDo2 and CTDo5 showed a 2-fold increase, whereas CTDa decreased (Fig. 6, A and D). When
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FIGURE 7. Presence of Tip110 at the HIV-1 LTR core promoter. A, schematic of the LTR core promoter and the primer locations for PCR amplification. B, Western blot analysis for Tip110 expression in 293T cells transfected with pNL4-3 and pTip110.His (293T*) or pcDNA3 (293T), U373MAGI containing the LTR promoter-driven β-galactosidase gene (MAGI), and CEM-GFP cells containing the LTR promoter-driven GFP gene (CEM.GFP). C and D, 293T cells transfected with the pNL4-3 and pTip110.His (C, bottom) or pcDNA3 (C, top), U373MAGI (D, top), and CEM-GFP cells (D, bottom) were subjected to the ChIP assay. Following chromatin cross-linking, shearing, and immunoprecipitation with α-CTDa, α-His, or α-Tip110 antibody, reverse cross-linking was performed, and the DNA was then purified and analyzed by PCR with the primer set specific for the HIV-1 LTR core promoter region, as shown in A. Input DNA and immunoprecipitated DNA without any antibody (No Ab) or with an isotype-matched IgG were included as the ChIP controls. Moreover, PCR with a primer set specific for a GAPDH coding region was performed and also included as the control. E and F, Jurkat cells were transfected with Tip110.His or pcDNA3 and then infected with NL4-3 (equivalent to 20,000 cpm of reverse transcriptase, which gave rise to about 80% infection in 3 days determined by intracellular p24 staining). Cells were harvested for Western blot analysis for Tip110 expression (E) or ChIP assay as described above (F).

cells were transfected with Tat alone, CTDa showed a slight decrease, and CTD05 exhibited little change, but CTD02 showed a 1.5-fold increase (Fig. 6, B and E). In comparison, when cells were transfected with both Tat and Tip110, CTD02 and CTD05 both increased by 3-fold, whereas CTDa decreased (Fig. 6, C and F). These results indicate that expression of both Tip110 and Tat led to enhanced phosphorylation of RNAPII at both serine 2 and serine 5, which may account for Tip110 function in transcription activation. Expression analysis of P-TEFb (cyclin T1 and CDK9) or TFIIH (cyclin H and CDK7) confirmed that Tip110, Tat, or both did not significantly alter the expression of these proteins (data not shown), indicating that increased RNAPII phosphorylation by Tip110 or Tat was not the result of any changes in the expression levels of P-TEFb or TFIIH.

To further ascertain the effects of Tip110 interaction with Tat and CTDa on RNAPII phosphorylation, we took advantage of the ΔNT mutant that did not bind to CTDa (Fig. 4) and Tat (32) and performed similar experiments. Compared with the cells that were transfected with Tat alone, expression of ΔNT did not show considerable changes in RNAPII phosphorylation (Fig. 6, G and H).

Tip110 Was Present on the HIV-1 LTR Core Promoter—To determine whether Tip110 was recruited to the HIV-1 LTR promoter in vivo, ChIP experiments were carried out. The HIV-1 LTR core promoter and the primers for the ChIP assay are illustrated in Fig. 7A. For the ChIP assay with exogenous Tip110, 293T cells were transfected with the proviral plasmids pNL4-3 and pTip110.His, and Tip110 expression was confirmed (marked by an asterisk; Fig. 7B). Chromatin from transfected cells was isolated and immunoprecipitated with anti-His antibodies followed by PCR to amplify the HIV-1 LTR core promoter region. ChIP with no antibody (No Ab) and an isotype-matched IgG was included as negative controls, whereas ChIP with anti-CTDa antibody was included as a positive control. Moreover, PCR with primers spanning the GAPDH coding region was also performed as the other specificity control. Furthermore, PCR with ChIP input DNA was performed to ensure the quality and quantity of the DNA. The LTR promoter was amplified from the anti-His (Tip110) immunocomplex of cells that were transfected with pTip110.His (Fig. 7C, bottom panels). As expected, the LTR promoter was amplified from the anti-CTDa immunocomplex of cells that were transfected with pcDNA3 or pTip110.His. These results indicate that exogenous Tip110 is present at the HIV-1 LTR promoter in vivo.

Considering the unintegrated nature of the HIV proviral DNA and overexpression of Tip110 in the above experimental settings, we proceeded to determine whether endogenous Tip110 was present in the integrated HIV-1 promoter. We took advantage of U373MAGI cells that have an integrated HIV-1 LTR promoter-driven LacZ transgene cassette and CEM-GFP cells that have an integrated HIV-1 LTR promoter-driven GFP transgene cassette. Western blot analysis was first performed to confirm expression of endogenous Tip110 in these two cell lines (Fig. 7B). Then chromatin was isolated from these cells, immunoprecipitated, and subjected to PCR as above. The LTR promoter was clearly amplified from both CTDa and Tip110 immunocomplex prepared from both CEM-GFP and U373MAGI cells (Fig. 7D). Moreover, human CD4+ T lymphocyte Jurkat cells were transfected with pcDNA3 or pTip110.His (Fig. 7E) and then infected with HIV-1 NL4-3 viruses. A similar ChIP assay was performed. The LTR promoter was amplified from Tip110 immunocomplex prepared from cells that were transfected with pcDNA3 or pTip110.His, whereas the LTR promoter was only amplified from His immunocomplex prepared
from pTip110. His-transfected but not pcDNA3-transfected cells (Fig. 7F). Collectively, these results suggest that endogenous Tip110 is also detectable at the integrated HIV-1 LTR core promoter in vivo.

Tip110 Expression Was Associated with P-TEFb Recruitment to the LTR Promoter and Phosphorylation of Serine 2 of the RNAPII CTD.—We then evaluated the relationship among Tip110 expression, P-TEFb recruitment to the LTR promoter, and RNAPII phosphorylation. 293T cells were first transfected with pLTR-Luc and pTat.Myc, pTip110.HA, or both. Then a ChIP assay was performed using immunoprecipitations against Myc (Tat), HA (Tip110), cyclin T1, CDK9, CTDa, CTDd2, or CTDd5, followed by PCR to amplify the HIV-1 LTR region (Fig. 8A). Tat and Tip110 protein, included as positive controls, could be detected at the LTR region. No CDK9 or cyclin T1 was detected at the LTR promoter in the absence of Tat, but in the presence of Tip110 overexpression, increases in the levels of cyclin T1 and CDK9 recruited to the LTR promoter were observed. Meanwhile, CTDd2 was only observed in the LTR promoter in the presence of Tat expression and exhibited a 2-fold increase in the presence of Tip110, whereas CTDd5 was detected without Tat expression and showed little change in the presence of Tip110 and Tat. As expected, neither P-TEFb nor RNAPII could be detected at the LTR promoter when the immunoprecipitation was performed using mouse IgG as a control.

We performed similar experiments in cells with down-modulated Tip110 expression using Tip110-specific shRNA. Tip110 siRNA expression led to drastic and lasting knockdown and subsequently cell death and was not used in these experiments (39, 40). Tip110-shRNA has been shown to reduce endogenous Tip110 expression at day 7, whereas its backbone vector pSIREN has no effect on Tip110 expression (see Fig. 9). We transfected Tip110-shRNA or its cognate control pSIREN into 293T cells at day 0, followed by transfection of plLTR-Luc and pTat.Myc at day 4. The cells were harvested at day 7 for ChIP analysis. We found that there was less CDK9 and cyclin T1 recruited to the LTR promoter, and only half of the CTDd2 was detected in the absence of Tip110 (Fig. 8B). Meanwhile, CTDa increased slightly, but CTDd5 decreased in the absence of Tip110. These data together suggest that Tip110 interaction with Tat enabled more P-TEFb to be recruited to the HIV-1 LTR promoter and further enhanced RNAPII serine 2 phosphorylation at the LTR promoter.

Tip110 and Tat Increased the LTR Transcription Elongation Efficiency.—To determine if Tip110 interaction with Tat had direct effects on HIV-1 transcription, we adapted an in vitro guanine (G)-less transcription/elongation assay (Fig. 9A) and allows synthesis of two RNaseT1-resistant transcripts: the promoter-proximal short transcript and the promoter-distal long transcript. Expression of these two transcripts has widely been used to determine transcription initiation and elongation, respectively (41). First, an in vitro transcription was set up to contain nuclear extract, the G-less DNA template, recombinant Tat, Tip110, or Tat and Tip110. After digestion of the template DNA with DNase I, synthesized RNA was recovered, and semi-quantitative RT-PCR was performed to determine the relative levels of the short and long G-less transcripts using primers (Fig. 9A) that were specific for each of those two transcripts expected to give rise to PCR products of 300 (ST) and 520 bp (LT), respectively. As expected, Tat increased long transcript expression but had little effect on short transcript expression (Fig. 9B). Compared with Tat, Tip110 also slightly increased expression of the long transcripts and had little effect on the expression of short transcripts. Compared with Tat or Tip110 alone, the presence of both Tat and Tip110 showed considerably enhanced expression of the long transcripts but had no effect on the expression of short transcripts. These results indicate that the interaction of Tip110 with Tat led to a higher efficiency of elongation for RNAPII complexes formed on the LTR promoter.

To further characterize the significance of Tip110 in Tat-mediated LTR transcription elongation, we knocked down endogenous Tip110 and determined its effects on the LTR transcription. To achieve this, we knocked down endogenous
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FIGURE 9. Direct effects on HIV-1 transcription elongation by Tip110 and Tat. A, schematic of the HIV-1 LTR double G-less cassette used in the experiment. The transcript derived from this cassette contains two G-less regions, a short upstream G-less region (ST) and a long downstream G-less region (LT), which have been used to determine transcription initiation and transcription elongation, respectively. The primer locations to amplify the short and long upstream G-less regions have been indicated. B, in vitro transcription assay was performed with linearized double G-less plasmid and 0, 0.1, and 0.2 μg of recombinant Tat protein or 0.2 and 0.4 μg of recombinant Tip110 protein or 0.1 μg of recombinant Tat protein plus 0.2 and 0.4 μg of recombinant Tip110 protein. After DNase treatment, RNA was isolated and RT-PCR-amplified using primers specific for the short and long upstream G-less regions. C and D, 293T cells were first transfected with pSIREN or psh-Tip110 at day 0 and then transfected with the double G-less cassette and pTat.Myc plasmid at day 4 and again at day 7. Cells were harvested on day 7 and day 10 for whole cell extracts, followed by Western blot analysis with Tip110, Myc, or α-β-actin antibody (C), or for RNA extraction, followed by RT-PCR analysis for the short and long upstream G-less region transcripts (D). Elongation efficiency was calculated as the ratio of long to short upstream G-less region transcripts. E and F, HeLa nuclear extract was first added to α-Tip110 antibody, an isotype-matched IgG, or no antibody (No Ab) and then protein A beads. After a 2-h incubation, the protein A beads were recovered by brief centrifugation and discarded, whereas the supernatant was saved and subjected to Western blot analysis for Tip110 (E) or used in the in vitro transcription assay in the presence of recombinant Tat or Tat plus Tip110, followed by RNA isolation and RT-PCR as described above (F). The data were representative of three independent experiments.

Tip110 in 293T cells using Tip110-specific shRNA and performed an in vivo HIVdG-less assay in these cells. We transfected pHIVdG-less plasmid with pTat.Myc into 293T cells on day 0, 4, or 7 following transfection with either Tip110-specific shRNA or the backbone vector pSIREN. Cells were harvested at day 7 or 10 for Western blot or semiquantitative RT-PCR analysis. Western blot analysis demonstrated that endogenous Tip110 protein expression was down-modulated by pshTip110 at day 7 post-transfection but showed some recovery at day 10, and Tip110 expression showed no significant changes with pSIREN transfection (Fig. 9C). The RT-PCR analysis showed that the long transcripts showed considerable and consistent increases in cells that were transfected with Tat and pSIREN (Fig. 9D, left). In contrast, in cells that were transfected with Tat and pshTip110, the long transcripts showed similar increases at day 0 when Tip110 expression was not affected, considerable decreases at day 7 when Tip110 expression was knocked down, and considerable recovery at day 10 when Tip110 expression returned (Fig. 9D, right). The apparent interdependent relationship provides additional evidence to support the important roles of Tip110 in HIV-1 LTR transcription. Taken together, these results further demonstrate the specific role of Tip110 in regulation of the RNAPII elongation efficiency on the LTR promoter.

Last, we determined if direct depletion of endogenous Tip110 would impede Tat-mediated LTR transcription. To this end, nuclear extract was immunodepleted of endogenous Tip110 using an anti-Tip110 antibody before tip was used in the in vitro HIVdG-less transcription assay as described above. Immunodepletion with an isotype-matched IgG and without any antibodies was also included as a control. Western blot analysis confirmed that immunodepletion with anti-Tip110 antibody removed more than half of the endogenous Tip110 from nuclear extract (Fig. 9E). The RT-PCR analysis showed that Tip110 depletion from nuclear extract led to a considerable decrease in synthesis of the long transcripts compared with the isotype-matched IgG-depleted nuclear extract (Fig. 9F). Interestingly, the addition of recombinant Tip110 back to the Tip110-depleted nuclear extract failed to restore synthesis of the long transcripts. The inability of recombinant Tip110 to restore the transcription elongation is probably due to removal of other transcription factors that are associated with Tip110 depletion, such as RNAPII itself.

DISCUSSION

Transcriptional activation of the HIV-1 LTR promoter is a complex event and requires the coordination of viral proteins...
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and several cellular proteins. One of the key factors involved in this activation is the Tat protein, which enhances HIV-1 transcription by promoting the formation of transactivation complexes at the LTR promoter via protein-protein interactions. In addition to Tat protein, there are a large number of cellular factors also involved in transcription. They either function by removing the negative inhibitor that blocks RNAPII phosphorylation at the promoter region or by recruiting the elongation-competent RNAPII-containing complex (21, 42, 43). In this study, we focused on the Tip110 protein (Tat-interacting protein of 110 kDa). Studies from our group have shown that Tip110 functions synergistically with Tat in Tat-mediated transactivation of the HIV-1 LTR promoter, thereby increasing viral gene expression and virus production (32).

To understand the underlying molecular mechanisms of Tip110, we first examined its interaction with transcription factors. A large number of cellular factors are involved in Tat-mediated transcription; many recruit elongation-competent RNAPII-containing complexes and have been shown to interact with multiple transcription factors. For example, SKIP, Tat-SF1, and CA150 are reported to associate with Tat-P-TEFb in nuclear extracts and are present in large RNAPII elongation complexes (21, 23, 44). Tip110 was first identified as a Tat-interacting protein from a yeast two-hybrid system of a human fetal brain cDNA library, using Tat as bait (32). We found that Tip110 bound to unphosphorylated RNAPII (CTDa) but not its phosphorylated form (CTDo) (Fig. 2). This direct and specific binding was further supported by the results of a GST pull-down assay (Fig. 3) and mutagenesis analysis (Fig. 4). Protein structure analysis predicted that the N-terminal two-thirds of the Tip110 protein contains seven HAT motifs. These HAT motifs provide a structural unit of two antiparallel α-helices that form functional TPR(s) and determine the specificity of protein-protein interactions (45). Besides binding to unphosphorylated RNAPII, this domain is also responsible for Tip110 interaction with Tat (32). This suggests that Tip110 forms a complex with Tat and unphosphorylated RNAPII through its N-terminal domain and may play a role in stabilizing the transactivation complex. In addition, the HAT domain of Tip110 interacts with a C-terminal region of the U4/U6 snRNP-specific 90-kDa protein that functions in the reassembly of the U4/U6 snRNP (26, 27). The HAT domain is also involved in Tip110 interaction with the splicing factor RNPS1 (46), suggesting that the HAT domain determines the specificity of Tip110 interactions with other proteins.

Although Tip110 and P-TEFb both co-localize in the nuclear speckle area (35), our data indicated that no interaction existed between them (Fig. 1). This phenomenon distinguishes Tip110 from other cellular transcription factors involved in the P-TEFb complex. For example, SKIP associates with P-TEFb and is recruited to the LTR promoter by Tat (21). ELL2 is another elongation factor in the P-TEFb complex, and Tat recruits more ELL2 to P-TEFb and assists in the stabilization of ELL2 to activate P-TEFb (47). Although there is no direct interaction between Tip110 and P-TEFb, Tip110 is capable of recruiting more P-TEFb to the transcription complex in the presence of Tat (Fig. 5). Therefore, we speculate that Tip110 might be first recruited to the unphosphorylated RNAPII on the LTR promoter in the absence of Tat and then assist in the recruitment of P-TEFb through Tat. After the RNAPII has been phosphorylated by P-TEFb and establishes transcription elongation, Tip110 would become disassociated from phosphorylated RNAPII and commence a new cycle of LTR transactivation.

Because P-TEFb is responsible for RNAPII serine 2 phosphorylation, we set out to investigate whether Tip110 or Tat would alter the RNAPII phosphorylation level. Our data showed that Tip110 protein alone could decrease the level of unphosphorylated RNAPII while increasing the level of both serine 2- and serine 5-phosphorylated RNAPII by 2-fold, suggesting that Tip110 is a weak transactivator of the HIV-1 LTR and leads to its basal level transcription. When Tip110 and Tat were co-expressed, an increase in serine 2 and serine 5 RNAPII phosphorylation levels coincided with a large decrease in unphosphorylated RNAPII (Fig. 6), which may account for the Tip110 function in Tat-mediated transcription. However, in these studies, we examined the overall RNAPII phosphorylation level in 293T cells in the presence of Tip110 and Tat, not the RNAPII specifically located on the LTR promoter. Several genes other than the HIV-1 LTR promoter are regulated at the stage of transcription elongation by P-TEFb, including hsp70 and proto-oncogenes c-myb, c-myc, and c-fos (48–50). For these genes, the RNAPII complexes are stalled in the 5’ region of the transcription unit, and P-TEFb recruitment is the key regulator that helps RNAPII to overcome this rate-limiting step. It is the TATA-box, not the TAR structure, that is important for the recruitment of P-TEFb (41). Therefore, the increased phosphorylated form of RNAPII may result from the recruitment of P-TEFb by Tip110 and Tat on these cellular genes. In order to examine RNAPII phosphorylation on the LTR promoter, one can label the 5’ end of the LTR-promoter templates with biotin and isolate LTR-bound preinitiation complexes to detect protein components associated with it (51). We would expect to see a greater increase in RNAPII phosphorylation located on the HIV-1 LTR promoter in the presence of Tip110 and Tat.

HIV-1 transcription is regulated by the interplay between a combination of viral and cellular transcription factors with binding sites located within the HIV-1 LTR promoter. Using the ChIP assay, we first detected Tip110 associated with the LTR promoter expressed transiently by transfecting pNL4-3 into 293T cells (Fig. 7). To further examine whether Tip110 could be recruited to the integrated LTR promoter in the absence of Tat, we performed a ChIP assay with endogenous Tip110 using U373MAGI or CEM-GFP cells. In both cell lines, the LTR promoter is integrated into the chromosome with various proto-oncogenes c-fos (48–50). For these genes, the RNAPII complexes are stalled in the 5’ region of the transcription unit, and P-TEFb recruitment is the key regulator that helps RNAPII to overcome this rate-limiting step. It is the TATA-box, not the TAR structure, that is important for the recruitment of P-TEFb (41). Therefore, the increased phosphorylated form of RNAPII may result from the recruitment of P-TEFb by Tip110 and Tat on these cellular genes. In order to examine RNAPII phosphorylation on the LTR promoter, one can label the 5’ end of the LTR-promoter templates with biotin and isolate LTR-bound preinitiation complexes to detect protein components associated with it (51). We would expect to see a greater increase in RNAPII phosphorylation located on the HIV-1 LTR promoter in the presence of Tip110 and Tat.
Tip110 recruits more cyclin T1 than CDK9 into the transcription complex, and this is most likely because Tat directly interacts with Tip110 and mediates the recruitment of cyclin T1 by Tip110. We further examined RNAPII phosphorylation levels at the LTR promoter by performing a ChIP assay with 8WG16, H5, and H14 antibodies. Serine 2 phosphorylation of RNAPII only took place in the presence of Tat and increased with Tip110 overexpression, possibly as a result of the increased recruitment of P-TEFb by Tip110 and Tat. Unphosphorylated and serine 5-phosphorylated RNAPII were readily detected at the HIV-1 promoter in the absence of Tat, whereas Tip110 supplied along with Tat produced a slight effect on serine 5 phosphorylation, consistent with previous reports that TFIIH is recruited to the LTR promoter for RNAPII serine 5 phosphorylation in the absence of Tat (38). The increased recruitment of T-PEFb to the LTR promoter and increased CTDo2 (Fig. 8) were associated with increased transcription elongation of the LTR promoter (Fig. 9) in the presence of Tip110, providing additional evidence that Tip110 is an important factor for HIV-1 Tat-mediated transcription.

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