RESEARCH ARTICLE

A novel non-radioactive assay for HIV-RT (RdDp) based on pyrosequencing for high-throughput drug screening

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ABSTRACT

Current in vitro assays for the activity of HIV-RT (reverse transcriptase) require radio-labeled or chemically modified nucleotides to detect reaction products. However, these assays are inherently end-point measurements and labor intensive. Here we describe a novel non-radioactive assay based on the principle of pyrosequencing coupledenzyme system to monitor the activity of HIV-RT by indirectly measuring the release of pyrophosphate (PP_i), which is generated during nascent strand synthesis. The results show that our assay could monitor HIV-RT activity with high sensitivity and is suitable for rapid highthroughput drug screening targeting anti-HIV therapies due to its high speed and convenience. Moreover, this assay can be used to measure primase activity in an easy and sensitive manner, which suggests that this novel approach could be wildly used to analyze the activity of PP_i-generated and ATP-free enzyme reactions.

KEYWORDS HIV-RT (RdDp), assay, non-radioactive, high throughput drug screening

INTRODUCTION

Human immunodeficiency virus (HIV) is a retrovirus that can lead to acquired immunodeficiency syndrome (AIDS), a condition in humans in which the immune system begins to fail, leading to life-threatening opportunistic infections. HIV type 1 (HIV-1) is a positive single-strand RNA virus and the genome encoded reverse transcriptase (RT) is essential for HIV-1 replication. During replication, the RNA-dependent DNA polymerase (RdDp) transcribes single-stranded RNA into double-stranded DNA. Normal transcription involves the synthesis of RNA from DNA; hence, reverse transcription is the reverse of this. The RT converts single-stranded viral RNA into a linear double-stranded DNA, which can be integrated into host chromosomes (Sarafianos et al., 2004). HIV-1 RT is a heterodimer consisting of a 66-kDa (p66) and a 51-kDa (p51) subunit. With the exception of the last 120 C-terminal residues in p66, which form the RNase H domain, these two polypeptide chains have the same 440 N-terminal residues and can be divided into thumb, palm, fingers and connection (Kohlstaedt et al., 1992; Jacobo-Molina et al., 1993).

Due to the essential role in the viral life cycle and clear functional and structural investigation, HIV-RT has been the target of considerable interest for anti-viral drug development to date. In previous studies, a number of assays had been developed and applied to monitor HIV-RT RdDp activity in vitro for drug screening. Initial HIV-RT assays were based on the measurement of generated radioactive production by using radioactive compounds as the substrates. The most conventional method is to use ³H-labeled dTTP for in vitro HIV-RT assays (Hizi et al., 1993; Antoun et al., 1994; Kanyara and Njagi, 2005). However, the conventional isotopic HIV-RT assay is labor intensive and is becoming increasingly restrictive due to the use of radioactive materials. Although radioactive signals are easily detected and quantified, these assay systems inherently produce end-point measurements and kinetic evaluations are not easy to carry out. Bearing in mind the radioisotopic hazard, non-radioisotopic assays for measuring RT activity using mainly colorimetric methods have

also been developed (Suzuki et al., 1993a, b). Briefly, BrdUTP is incorporated into a DNA strand complementary to the polyA template. Subsequently, an anti-BrdU antibody conjugated to alkaline phosphatase is added and the amount of incorporated BrdU detected using a substrate pNPP. The color intensity of each well is read using a standard plate reader (405 nm, reference at 620 nm) (Vassiliou et al., 2000). Several assays have been also established based on similar principles, but with variation in nucleotides modification and color developing methods (Porstmann et al., 1991; Urabe et al., 1992; Suzuki et al., 1993a, b; Odawara et al., 2002). Product-enhanced reverse transcriptase (PERT) assays were developed in order to increase the sensitivity. The principle of the PERT assay involves producing cDNA from template RNA by suspected RT in a test sample, amplifying the cDNA by polymerase chain reaction (PCR) and detecting the PCR amplicons by gel electrophoresis (Chang et al., 1997), Southern blotting (Silver et al., 1993), or an enzymelinked oligonucleotide sorbent assay (ELOSA) (Pyra et al., 1994; Andre et al., 2000). To further increase its sensitivity, the fluorescence PERT (F-PERT) assay and single-tube fluorescent product-enhanced reverse transcriptase (STF-PERT) assay have been improved using real-time RT-PCR, with real-time detection of PCR product levels during amplification by using TagMans PCR technology (Arnold et al., 1998; Lovatt et al., 1999). Though RT-PCR amplification F-PERT and STF-PERT assay could be real-time and high sensitive, a real-time thermal cycler is still required and the speed of this assay could not perfectly fit high-throughput drug screening.

Recently, a new sensitive and continuous non-radioactive RdRp assay in which activity is measured based on the detection of pyrophosphate (PP_i) has been reported (Lahser and Malcolm, 2004). Free PP_i is released when the nucleotide is incorporated onto the provided template. The PP_i can then be detected by a coupled-enzyme system which can convert PP_i to ATP and finally to a light signal. In comparison with the other non-radioactive assays described above, this assay has a competitive advantage in that native NTPs can be used. The coupled-enzyme system greatly amplifies the signal, thus allowing for the use of lower concentrations of enzyme, which is important in high-throughput inhibitor screening aiming at polymerase. Meanwhile, data in this assay are collected in real time, which makes the kinetic analysis of the whole reaction easy to assess.

Considering the similarity between RdRp and RdDp, we applied the same approach to an HIV-1 RT *in vitro* activity assay. Our results show that this high-throughput method can measure HIV-RT activity easily and in real-time, suggesting that it is an effective assay and could potentially have significant impacts in high-throughput drug screening. Moreover, this assay could be used to measure primase activity easily and sensitively, which implies that this assay could be wildly used to monitor the activity of PP_i-generated and ATP-free enzyme reactions.

RESULTS

Enzyme concentration

The reaction was performed as indicated in "Materials and Methods". Signals were collected every 10 s with an integrated time of 100 ms, then the real-time light intensity was recorded (Fig. 1A). As the assay can be considered as a pseudo-first order reaction, which can be described by $P = S_0(1 - e^{-kt})$, the initial stage of the reaction should simply follow the Taylor expansion $P = S_0kt$. We therefore used linear regression to determine the velocity. Data used for



Figure 1. Effects of a series of HIV-RT concentrations. (A) Titration of HIV-RT. HIV-RT was serially diluted in storage buffer and assayed (n = 5) at room temperature with 2 µM dTTP, 15 mU ATP sulfurylase, and the poly A/oligo (dT) 18 in a 96-well plate with a final reaction volume of 100 mL. Light was captured for 100 ms every 30 s. Initial velocity values were derived from progress curve analyses, fitted by linear regression to the Taylor expansion of a simple first order model: $P = S_0 kt$, where we apply $S_0 k$ to evaluate the relative initial velocity. (B) Linearity of the reaction with HIV-RT concentration. Relative initial velocity estimates from the data in (A) were plotted against [HIV-RT]. Values were fitted by linear regression.

initial velocity determination were corrected with a non-HIV-RT control, and the reaction rate was linear with respect to enzyme concentration in the range tested (Fig. 1B).

dTTP concentration

The effects of dTTP concentration were evaluated in the reaction system. As indicated by Fig. 2A, the initial velocity was linear with respect to the concentration of dTTP in the range $2-10 \,\mu$ M (Fig. 2B). A little inaccuracy was observed at low concentration and will be discussed in further detail below.

Oligo (dT) 18 concentration

As the primer of the first reaction, the effects of oligo (dT) 18 concentration were also investigated (Fig. 3A). Polymerase activity increased with primer concentration, with indications of saturation at the maximum concentration tested (2 μ g/mL) (Fig. 3B).

Pre-incubation

The impact of pre-incubation of HIV-RT with template (primer) was examined using the assay described above. HIV-RT was co-incubated with template (primer) prior to the initiation of RNA synthesis by addition of the remaining reaction components. As shown in Fig. 4, there is no obvious increase in signal within 3 h of pre-incubation. The results suggest that, for this system, formation of the initiation complex was not rate limiting, indicating that a good signal can be achieved without pre-incubation.

Inhibition of HIV RT

To demonstrate the utility of this assay for the evaluation of inhibitor, the commercial inhibitor nevirapine was applied to the assay. Different doses of nevirapine were introduced to the assay and compared with positive (HIV-RT+) and negative (HIV-RT-) controls. Signals were collected and initial



Figure 2. Effects of dTTP concentration on activity. (A) Various concentrations of dTTP were added; the HIV-RT concentration is 1.25 μ g/mL; other conditions and parameters are as described for (Fig. 1A). Results shown are the average of 5 independent experiments (*n* = 5). (B) Linearity of the reaction with dTTP concentration. Relative initial velocity estimates from data in (A) were plotted against [dTTP].



Figure 3. Effects of oligo (dT) 18 concentration on activity. (A) Various concentrations of oligo (dT) 18 were added; the HIV-RT concentration is 1.25 mg/mL, other conditions and parameters are as described for Fig. 1A. Results shown are the average of 5 independent experiments (n = 5). (B) Polymerase activity increased with primer concentration, with indications of saturation at the maximum tested (2 µg/mL).



Figure 4. Pre-incubation of HIV-RT. HIV-RT was preincubated with polyA and oligo (dT) 18; the HIV-RT concentration is 1.25 mg/mL, other conditions and parameters are as described for Fig. 1A. No obvious increase in signal was observed within 3 h of preincubation. Data were collected from three independent reactions (n = 3) for 0.1 s every 30 s during a 6-min reaction.

velocities were calculated to evaluate the specific activity. Dose-dependent reduction in signal was observed for the nevirapine (Fig. 5).



Figure 5. Inhibitor studies using nervirapine. (A) Inhibition of HIV-RT with varying concentrations of inhibitor nevirapine compare with positive (HIV-RT+) and negative (HIV-RT-) controls (n = 3). Reactions contained HIV-RT (1.25 µg/mL); other conditions and parameters are the same as described for Fig. 1A. Luminescence was measured for 0.1 s every 60 s during a 15-min reaction.

Comparison with colorimetric assay in sensitivity

Analysis of a serial dilution of HIV-RT in low concentrations was done in comparison with a well accepted colorimetric assay to determine assay sensitivity. In this comparison, the same serial dilutions of recombinant HIV-RT were measured. As shown in Table 1, the detective sensitivity of PP_i enzyme

coupled assay is comparable with colorimetric assay in our experimental condition. The colorimetric assay can detect as low as 30 ng/mL recombinant HIV-RT after 90-min reaction time, while our data indicate PPi coupled enzyme assay can reach the equivalent sensitivity after only 15-min reaction time.

DISCUSSION

As mentioned above, the reliability of the assay relies in the fact that the components in the assay have to be carefully adjusted to ensure that the PP_i generating reaction is the ratelimiting step. First, the substrate and enzyme in the light-generating cascade, such as APS, D-luciferin, ATP-sulfrylase and luciferase, must be sufficient to ensure that all the PP_i generated by the HIV-RT can be converted into a light signal.

Second, the substrates required for PP_i generation, such as polyA, oligo (dT) 18 and dTTP, should be in excess to ensure their concentrations remain nearly constant at the beginning of data collection, so that the initial velocity is solely dependent on the specific activity of HIV-RT. According to our experiments, the linear relationship between HIV-RT activity and initial velocity can be guaranteed at the concentrations of HIV-RT ($0-5 \mu g/mL$) and dTTP ($1-5 \mu M$) tested, when sufficient oligo (dT) 18 is provided ($1 \mu g/mL$).

Signal-to-noise ratio is another issue that must be taken into consideration in this assay. As shown above, there is a little inaccuracy when the signal is collected at low dTTP concentration. The signal-to-noise ratio is not as good compared with that at high light intensity. To improve the signal-to-noise ratio at low concentration, the integration time can simply be extended to 0.2 s in order to enhance the light intensity. On the other hand, both the concentrations of HIV-RT and dTTP can be increased properly to overcome this problem, but the velocity of the PP_i generating reaction has to be limited to a range in which all of the accumulated PP_i can be quickly converted to a light signal.

Comparing with the other assays such as F-PERT that has a better sensitivity, the PP_i-coupled enzyme assay has the advantage in convenience and time saving. Moreover, the sensitivity can be improved by simply increase the amount of the components if needed.

Our data clearly show that this assay is able to demonstrate the inhibition by nevirapine, which is a non-competitive inhibitor of HIV-RT. Nucleoside inhibitors of RT, i.e., the triphosphates of AZT or ddC, will release PP_i as they are added in the chain termination reaction. Dose- and time-dependent reduction in signal was observed for the compound (Lahser and Malcolm, 2004), which is consistent with chain termination (i.e., once elongation is terminated by 3'-ddTTP incorporation, no further PP_i can be produced). However, considering that we apply homopolymer (PolyA) rather than heteropolymer as a template, only deoxythymidine analogs (ddTTP, AZTTP etc.) can be evaluated. A special heteropolymer should be synthesized as a template to make better evaluation of all kinds of nucleoside inhibitors.

Table 1	Comparison of	colorimetric assay	and PP _i	coupled	enzyme	assay
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HIV RT (μg/mL)	relative value				
	colorimetric assay (absorbance 450 nm)	PP _i coupled enzyme assay (light intensity)			
	90 min	15 min	30 min		
1	2.46325 ^a	NT ^b	NT		
0.1	0.98075	2300.052 ^c	4435.39		
0.03	0.4925	886.892	1505.82		
0.01	0.2475	496.812	662.996		
0.003	0.1895	444.566	539.364		
0.001	0.17575	333.866	374.84		
0	0.16175	276.398	366.984		
cutoff value ^d	0.33	553	734		

^a The result for colorimetric assay is based upon quadruplicate dilutions.

^b Not tested.

 $^{\rm c}$ The result for ${\sf PP}_{\rm i}$ coupled enzyme assay is based upon quintuplicate dilutions.

^d Cutoff value = 2×negative control.

In contrast to a similar assay adapted to monitor activity of the hepatitis C virus nonstructural protein 5B (NS5B) RdRp (Lahser and Malcolm, 2004), our observations indicate that pre-incubation does not give rise to a significant increase in the signal. This would suggest that, for HIV-RT, formation of the initiation complex was not rate limiting, and that a good signal can be achieved without pre-incubation. Therefore, this is a robust time-saving method that can easily be adopted for high throughput screening.

Moreover, some other results show that this enzyme assay can be also easily applied to primase activity measurement (unpublished data), which just excluded primer from this assay and select suitable template and substrate. This result suggests that this pyro-sequencing-based enzyme assay could be not only applied to high-throughput drug screening aiming at HIV-RT, but also could be wildly used in investigations of ATP-free and PP_i-produced enzyme systems.

In summary, the coupled enzyme system used in pyrosequencing was adapted to evaluate the specific activity of HIV-RT. The reaction system can provide reliable initial velocities at various enzyme and substrate concentrations in as little as 4 min compared to the assays described previously. This method can be easily performed in 96-well plates and signals can be rapidly collected in real time. These advantages ensure that the assay meets the demands of high-throughput anti-HIV-RT inhibitor screening.

MATERIALS AND METHODS

Principle

Pyro-sequencing is a method of DNA sequencing (determining the order of nucleotides in DNA) based on the "sequencing by synthesis" principle, which was innitially described by Ronaghi et al. (1998). This method relies on detecting the activity of DNA polymerase with a

chemiluminescent enzyme. The single strand DNA (ssDNA) template is hybridized to a sequencing primer and incubated with four enzymes (DNA polymerase, ATP sulfurylase, luciferase and apyrase), and the substrates adenosine 5'-phosphosulfate (APS) and D-luciferin. Once the additional dNTPs (ATP α S instead of ATP) are added into this system, the DNA polymerase incorporates the correct, complementary dNTPs onto the template and releases PP_i from each base pair. In the presence of APS, ATP sulfurylase coverts the released PP_i to ATP, which will help luciferase convert D-luciferin to oxyluciferin. This light, which was generated by oxyluciferin, will then be monitored by camera. The reaction steps proceed as follows:

$$(\text{DNA})_n + \text{dNTP} \xrightarrow[\text{(slow)}]{} (\text{DNA})_{n+1} + \text{PPi},$$
 (1)

$$APS + PPi \xrightarrow{(fast)} ATP + SO_4, \qquad (2)$$

and

 $\label{eq:D-luciferin} \begin{array}{c} \text{D-luciferin} + \text{ATP} + \text{O}_2 & \xrightarrow[luciferase]{} \text{vsyluciferin} + \text{AMP} + \text{PPi} + \text{CO}_2 & (3) \end{array}$

As the net result of the last two steps does not consume any PP_i produced by DNA polymerase, the polymerase activity can be easily evaluated by the PP_i accumulation in the first step. On the other hand, from the last equation, it is evident that the light intensity is proportional to the PP_i concentration in the reaction system. Thus the polymerase activity can be reasonable evaluated by the increase of light intensity.

As HIV-RT is a type of RdDp, it opens up the possibility that this system can be applied to monitor HIV-RT activity. In this system, HIV-RT, polyA, oligodT and dTTP were respectively used as DNA polymerase, template, primer and dNTP in the pyrosequencing system. Since the template, primer and nucleotide are consistent in all reactions, the light signal increase should be only related with HIV-RT activity in a unit time.

Materials

Chemical reagents for assay-related buffers were obtained from Sigma (St. Louis, MO). Purified firefly luciferase and D-luciferin were

from Sigma. dTTP was purchased from Takara Bio Inc. Adenosine 5phosphosulfate (APS) and adenosine 5-triphosphate (ATP) sulfurylase (1 U produces 1 µmol of ATP from APS and PP_i per min at pH 8.0 at 30°C) were obtained from Sigma. The polyA homopolymer was from Sigma, and the oligo (dT) 18 was synthesized from Invitrogen. Nevirapine was a gift from Prof. Yiming Shao from the China Center for Disease Control and Prevention (CDC). White flat bottom polystyrene microplates (96-well) for the polymerase/luciferase assay were from Corning (Corning Co., USA). A microtiter plate luminometer (Luminoskan Ascent) was obtained from Thermo Scientific.

The express construct of HIV-1 RT was given by Prof. Eddy Arnold as a kindly gift. The protein of HIV-1RT was expressed and purified following the reported procedure (Clark et al., 1995). Briefly, the cells were sonicated by using lysis buffer, which contains 50 mM sodium phosphate (pH 8.0), 600 mM NaCl, 0.1% Triton X-100, 5 mM BME, 5 mM Imidazole and 1 mM PMSF. After being centrifuged at 18,000 rpm 30 min, the supernatant was added into Ni-NTA column and target protein was eluted by 50 mM sodium phosphate (pH 8.0), 200 mM NaCl, 5 mM BME, 250 mM imidazole (pH 8.0). Subsequently, the ion-exchange column MonoQ was used to purify target protein. Enzyme stocks of the HIV-RT were prepared and stored in 10 mM Tris-HCl (pH 8.0), 75 mM NaCl.

Methods

Polymerase assay

Polymerase assays were performed at room temperature using a 100 μL reaction mix in a 96-well plate. Final reaction conditions for HIV-RT were 20 mM Hepes (pH 7.5), 10 mM MgCl₂, 60 mM NaCl, 1 μ g/mL oligo (dT) 18, 5 μ g/mL polyA, 2 μ M dTTP, and HIV-RT. The enzyme/template mixtures were supplemented with components for the ATP sulfurylase and luciferase coupled-enzyme reactions (310 μ M D-luciferin, 1 nM luciferase, 10 μ M APS, 15 mU ATP sulfurylase).

Ninty-six-well microtiter plates containing 50 μ L initiated enzyme/ template mix were supplemented with substrate mix by multi-channel pipette, then the plate was immediately transferred to the luminometer (Thermo Scientific) for detection of the light signal in real time (using the instrument's "kinetic" programme). A 0.1 s reading of each well was taken every 10 s (at least 20 s is needed if all the 96 wells are tested). All of the light signals collected were compared with corresponding non-HIV-RT controls. Data used for initial velocity calculations were restricted to the first 200 s to avoid the influence of substrate depletion and/or product inhibition.

The key point of the assay is to ensure that the generation of PP_i, i.e., the first reaction, is rate limiting. To ensure that, both enzymes and substrates should be carefully evaluated during the reaction. According to the above principle, the initial velocity should depend completely on the first reaction when the components in the light generating cascade are fixed and sufficient. Furthermore, the substrates in the PP_i generating reaction must be in excess to ensure linearity at the beginning of data collection. Thus, the effects of both the enzyme and substrates in the first step reaction were evaluated so that proper concentrations can be determined to guarantee the linear relation between HIV-RT activity and light intensity.

To determine whether the reaction system was reliable for evaluation of the specific activity of HIV-RT, a series of enzyme concentrations were applied to monitor the relationship between the specific activity and the initial velocity. To test the effects of dTTP on the initial velocity, various dTTP diluents were introduced to the assay. Reactions were performed as described above, with different dTTP concentration in the substrate mixture.

To assess the effects of the oligo (dT) 18 on the initial velocity so that a proper concentration can be determined, a series of oligo (dT) 18 concentrations were used to get a saturated concentration.

The effects of pre-incubation were assessed by combining (at 0–3 h) HIV-RT, polyA, and oligo (dT) 18 at twice the final reaction concentration and initiated by addition of an equal volume of a 2 \times reaction mix containing luciferase, ATP sulfurylase, dTTP, APS, and D-luciferin. Data were collected for 6 min.

To demonstrate the utility of this assay for the evaluation of HIV-RT inhibitors, it was applied to the commercial inhibitor nevirapine.

Comparison with colorimetric assay in sensitivity

To further demonstrate the validity and the sensitivity of the assay, analysis of a dilution series of HIV RT was done compare with a widely accepted colorimetric RT assay (Urabe et al., 1992).

Poly A-linked colorimetric RT assay was performed using the method described before (Suzuki et al., 1995) with slight modifications. Briefly, wells were prepared prior to the assay by $50 \ \mu$ L of 1.875 µg/mL poly A (Sigma) in 0.95 mM N-hydroxysulfosuccinimide (NHSS) (Shanghai Medpep Co., Ltd.) to 50 µL of 3.0 mM 1-ethyl-3-(3dimethylaminopropyl) carbodiimide (EDC) (Shanghai Medpep Co., Ltd.) per microtiter well (Covalink from Nunc), incubating for 14-18 h at room temperature and then storing at 4°C. Wells were washed 3 times with washing buffer (10 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 mM EDTA, 0.01% Tween-20) just prior to use. Enzyme activity was measured by adding 10 µL RT to 50 µL of reaction buffer (50 mM Tris-HCl, pH 8.0, 6 mM MgCl₂, 75 mM KCl, 5 mM DTT, 1 mg/mL BSA, 0.3125 µg/mL oligo (dT) 15 (Invitrogen), 0.52 µM dTTP (Takara), 0.26 µM Biotin-11-dUTP (Shanghai Mol-Immuno Biochemistry Lab)), in microtiter wells. The reactions were incubated at room temperature for 1.5 h and then washed with wash buffer. After washing, 100 µL of streptavidin phosphatase (SAP) (KPL[™]) 0.5 mg/mL diluted 1:1000 in PBS buffer with 1% BSA was added to each well. The plate was incubated for 30 min at 37°C. Free conjugate was removed by washing 6 times with washing buffer. As a final step, 200 µL 4-nitrophenyl phosphate disodium salt hexahydrate (pNPP) (Fluka) 1 mM in pNPP buffer (1 mM MgCl₂, 100 mM Bicine, pH 9.8) was added to each well, the plate was incubated at room temperature for 5-10 min and then read at 450 nm with a plate reader (Dynatech).

PP_i coupled enzyme assay was performed as described before.

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ABBREVIATIONS

APS, adenosine 5'-phosphosulfate; ATP, adenosine 5-triphosphate; BrdUTP, bromodeoxyuridine triphosphate; ELOSA, enzyme-linked oligonucleotide sorbent assay; F-PERT, fluorescence PERT; HIV-RT, human immunodeficiency virus reverse transcriptase; PERT, product enhanced reverse transcriptase; pNPP, p-nitrophenyl phosphate; PP_i, pyrophosphate; RdDp, RNA-dependent DNA polymerase; SA-ALP, streptavidin-alkaline phosphatase; STF-PERT, single-tube fluorescent PERT

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