FRET-based assay to screen inhibitors of HIV-1 reverse transcriptase and nucleocapsid protein

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ABSTRACT

During HIV-1 reverse transcription, the single-stranded RNA genome is converted into proviral double stranded DNA by Reverse Transcriptase (RT) within a reverse transcription complex composed of the genomic RNA and a number of HIV-1 encoded proteins, including the nucleocapsid protein NCp7. Here, we developed a one-step and one-pot RT polymerization assay. In this in vitro assay, RT polymerization is monitored in real-time by Förster resonance energy transfer (FRET) using a commercially available doubly-labeled primer/template DNA. The assay can monitor and quantify RT polymerization activity as well as its promotion by NCp7. Z-factor values as high as 0.89 were obtained, indicating that the assay is suitable for high-throughput drug screening. Using Nevirapine and AZT as prototypical RT inhibitors, reliable IC₅₀ values were obtained from the changes in the RT polymerization kinetics. Interestingly, the assay can also detect NCp7 inhibitors, making it suitable for high-throughput screening of drugs targeting RT, NCp7 or simultaneously, both proteins.

INTRODUCTION

During HIV-1 reverse transcription, the reverse transcriptase (RT) transcribes (+) strand viral RNA genome to form integration-competent double stranded DNA (1). RT is a p66/p51 heterodimer that exhibits RNA- and DNA-directed DNA polymerase and RNAse H activities. The p66 subunit is enzymatically active and can be structurally divided into thumb, palm, fingers and connection subdomains and RNase H domain (2,3). Although the p51 subunit shares the same polypeptide chain as the p66 subunit with the exception of the last 120 C-terminal amino acids, its role seems to be purely structural (2–4). Because of its critical role in reverse transcription and thus in retroviral replication, RT is an important target for anti-HIV drugs and two well represented classes of approved RT-inhibitor drugs are available (5–7). The first class called nucleoside reverse transcriptase inhibitors (NRTI) consists of nucleotide analogues such as Zidovudine (AZT), which directly compete with dNTP monomers to block the extension of the primer. The second class called non-nucleoside reverse transcriptase inhibitors (NNRTI) is composed of molecules structurally unrelated to nucleotides that bind non-competitively to RT allosteric sites (8,9). Both of these inhibitor classes, substantially slow down the progression of the virus. However, even when the two classes are used in association, the high prevalence of mutations in the HIV-1 genome results in the selection of drug resistant strains, leading to frequent viral escapes over the long term patient’s therapeutic management. In this context, a continuous development of new RT inhibitor drugs targeting the enzymatic activities of RT is needed (10–12). Alternatively, since RT is regulated by a number of other HIV-1 proteins in the reverse transcription complex (RTC), these proteins may constitute additional targets to block the reverse transcription process. For instance, an attractive complementary target could be the HIV-1 nucleocapsid protein NCp7 (13–15), a highly conserved 55-amino acid protein, characterized by two zinc finger motifs, that is quantitatively the main HIV-1 protein in RTC. NCp7 is thought to promote reverse transcription through its nucleic acid chaperone activity (16–21), but also by stabilizing the complex of RT with its substrate (22–24) and/or through direct interaction with RT (25–27).

Multiple assays have been reported to investigate and monitor RT polymerization activity. Most of these assays are usually discontinuous multistep assays, using radioactively or fluorescently labeled nucleotide analogs (28–42) and requiring extensive sample handling. Recently, a number of fluorescence-based approaches have been developed to further characterize the kinetics and mechanism of RT (43–47). Based on these approaches, RT was notably shown to dynamically flip between two opposite orientations on its substrate, namely a polymerase conformation, where the finger sub-domain is toward the primer 3’ end and a RNAse...
H conformation, where the RNase H domain is toward the primer 3' end, explaining how the binding mode of RT tunes its enzymatic activity (43,46). Based on these data, we recently developed a one-step and one-pot assay using Förster resonance energy transfer (FRET) between fluorescently labeled RT and primer/template (p/t) that allows monitoring RT flipping and polymerization in real time (48). This assay was shown to discriminate non-nucleoside RT inhibitors from nucleoside RT inhibitors and to determine reliably their potency. Though this assay is highly sensitive and convenient, it suffers from the need to produce a RT mutant with a single accessible Cys, to label it by a fluorophore and to purify it, in an enzymatically active state. Therefore, this assay can hardly be used with wild-type RT or with RT mutants resistant to RT inhibitors. In order to overcome these limitations, we proposed here a continuous one-pot FRET-based polymerization assay, using a commercially available doubly-labeled p/t similar to that previously used in single molecule experiments for monitoring RT polymerase activity (46). Through this assay, the RT-mediated extension of the primer as well its promotion by NCP7 could be monitored in real time, by conventional fluorescence techniques. Therefore, we propose here a simple and robust assay for screening and validating anti-RT and anti-NCP7 molecules.

MATERIALS AND METHODS

Proteins

Recombinant, heterodimeric mutant of HIV-1<sub>HRH10</sub> RT was expressed in Escherichia coli and purified as described previously (49). The expression system and purification protocol allowed the preparation of large quantities of heterodimeric enzyme in a homogeneous form. The prepared RT contains a E478Q mutation in its RNase H domain, but it was previously checked that this mutation does not affect the polymerase activity (50). RT labeled in its p51 thumb subdomain by Alexa 488 was prepared and its activity was tested as described (48). Proteins were stored at −80°C. Protein concentrations were determined using extinction coefficients at 280 nm of 260 450 M<sup>−1</sup> cm<sup>−1</sup>.

Full length NCP7 peptide was synthesized by solid-phase peptide synthesis on a 433A synthesizer (ABI, Foster City, CA, USA) as described (51). Purification by HPLC was carried out on a C8 column (Uptisphere 300 A, 5 μm; 250 × 10, Interchim, France) in 0.05% trifluoroacetic acid (TFA) with a linear gradient of 10–70% of acetonitrile for 30 min. The peptide purity and molecular weight were checked by LC/MS. An absorption coefficient of 5700 M<sup>−1</sup> cm<sup>−1</sup> at 280 nm was used to determine its concentration.

Nucleotides, RT-inhibitors

The deoxynucleotide (dNTP) mixture was prepared by mixing each of UltraPure dATP, dCTP, dGTP and dTTP, purchased from Sigma Aldrich. Nevirapine (11-Cyclopentyl-3,11-dihydro-4-methyl-6H-dipyrido[3,2-b:2',3'-e][1,4]diazepin-6-one) was also purchased from Sigma Aldrich and its stock solution was dissolved in dimethyl sulfoxide (DMSO). AZT triphosphate (5'-Azido-3'-Deoxythymidine-5 Triphosphate) was purchased from Genexxon bioscience (Germany).

Oligonucleotides

Synthetic purified oligodeoxynucleotides were purchased from IBA (Göttingen, Germany) with the necessary modifications and their concentrations were determined by UV absorbance at 260 nm by using the extinction coefficients provided by the supplier. FRET measurements were carried out using a 23-mer (5'- CAG CAG TAC AAA TGG CAG TAT TC) DNA-primer labeled at the T19 position with Cyanine 5 (Cy5), annealed to a 63-mer (3'-TGT CGT CAT GTT TAC CGT CAT AAG TAG GTG TTA CTA GTC CGA TTT CCC CTA GTC CGA CCC ATG)-template labeled at the T2 position with carboxytetramethylrhodamine (TMR). Both TMR and Cy5 were covalently attached by supplier, via a C6 amino link to their respective T residues in the primer and the template. Primer and template oligodeoxynucleotides were annealed by heating equimolar amounts in buffer at 90°C for 2 min, followed by cooling to room temperature over several hours in a heating block. Unless noted otherwise, all experiments were routinely carried out at 20°C in a buffer containing 50 mM Tris–HCl (pH 7.5), 10 mM KCl and 6 mM MgCl<sub>2</sub>.

Fluorescence measurements

All steady-state and kinetic experiments were performed using the doubly-labeled p/t duplexes, which are mentioned above, in the presence of HIV-1 RT. All measurements were performed at 20°C using either a Fluoromax 4 spectrofluorometer (Jobin-Yvon Horiba) or a stopped-flow apparatus (SF3, Biologics). The FRET donor, TMR, was excited at 540 nm, and its emission was recorded at 580 nm. Nucleotide incorporation kinetics was triggered by addition of deoxiribonucleotides (dNTPs) in excess to a pre-incubated mixture of HIV-RT and p/t duplexes at equimolar concentrations. The kinetics was fast enough to monitor the fluorescence intensities continuously without photobleaching. The annealing kinetic traces were adequately fitted using:

\[
I(t) = I_F - (I_F - I_0)(ae^{-k_{obs1}(t-r_0)} - (1-a)e^{-k_{obs2}(t-r_0)})
\]

where \( I_0 \) is the dead time, \( k_{obs1,2} \) are the observed kinetic rate constants, \( a \) is the amplitude of the fast component, and \( I_F \) and \( I_F \) are the fluorescence intensities before dNTPs addition and at completion of the reaction, respectively. The \( I_0 \) value was obtained from the steady-state fluorescence spectrum of the doubly labeled p/t duplex in the presence of RT and was thus fixed. All fitting procedures were carried out with Origin™ 8.6 software using nonlinear, least-square methods and the Levenberg-Marquardt algorithm. Analytical analysis was systematically cross-validated using Bayesian inference with Markov Chain Monte Carlo and Gibbs sampling (see Supplementary Data and Supplementary Figures S1 and 2).

Measurements in high-throughput screening (HTS) format

Experiments were performed in 96-well plates. The complex of RT with p/t was formed by mixing 100 nM RT and 100 nM doubly-labeled p/t duplex in a total volume of 200 μl. All measurements were performed at 20°C using a FLX-Xenius plate reader (Safas Monaco). Excitation wavelength
was 540 nm and emission was recorded at 580 nm using 10 nm slits. Polymerization kinetics were triggered by addition of 100 μM dNTPs. When kinetics were recorded in the presence of RT-inhibitors, the RT–p/t complex was pre-incubated with inhibitors for at least 5 min, unless otherwise stated. Each set of experiments was performed in three independent triplicates.

RESULTS AND DISCUSSION
Design and validation of the assay
A HIV-1 relevant DNA/DNA p/t sequence was selected from the HIV-1 pol gene. This p/t sequence lies outside any reported pausing-site (52) and thus, can be used to mimic a naturally occurring polymerization step during the plus strand synthesis of the HIV-1 genome. The doubly labeled p/t duplex was prepared by heat-annealing a 63-nt long DNA template labeled close to its 5' end at its T2 position by TMR, used as a FRET donor, with a 23-nucleotide long DNA primer labeled at its T19 position with Cy5, used as a FRET acceptor (Figure 1A). The TMR label was placed 38 nt away from the 3' end of the primer and 42 nt away from the Cy5 label and thus should not perturb RT binding.

Heat-annealed duplexes of labeled primer and template resulted in an FRET efficiency of 65% (Figure 1B, inset). This high value indicates that the average inter-dye distance is very short (~4.5 nm), suggesting that the single-stranded domain of the template is highly flexible, bringing the 5' end of the template close to the 3' end of the unextended primer (Figure 1A). Formation of the complex between RT and DNA p/t (hereinafter referred to as E.DNA) led to a marginal decrease in FRET (from 65 to 63%), indicating that RT binding induces marginal changes in the distance between the two chromophores. In contrast, addition of 100 μM dNTPs led to a progressive increase in TMR fluorescence that reached a plateau, corresponding to 16% FRET, in about 45 min (Figure 1B), due to the increase in the distance (from ~4.5 to 7 nm) between the fluorescent dyes as the full ds-duplex forms. As a control, neither the binding of RT nor the extension of the duplex was found to significantly affect the fluorescence of the same duplex labeled only with TMR (Supplementary Figure S3), clearly indicating that the FRET changes observed in Figure 1 were reflecting the polymerization activity of RT.

The progress curve in Figure 1B could be adequately fitted by a biexponential equation (Equation 1) with $k_{obs1} = 19 \pm 1 \times 10^{-3}$ s⁻¹, $k_{obs2} = 17 \pm 3 \times 10^{-4}$ s⁻¹ and $a = 0.35 \pm 0.1$. Fitting of the red trace according to Equation (1) provides $k_{obs1} = 14 \pm 1 \times 10^{-3}$ s⁻¹, $k_{obs2} = 13 \pm 3 \times 10^{-4}$ s⁻¹ and $a = 0.85 \pm 0.1$ and fitting of the blue trace provides $k_{obs1} = 18 \pm 2 \times 10^{-2}$ s⁻¹, $k_{obs2} = 63 \pm 5 \times 10^{-4}$ s⁻¹ and $a = 0.5 \pm 0.1$. Excitation and emission wavelengths were 480 and 520 nm, respectively. (C) Polymerization kinetic trace of 100 nM E.DNA (gray), 100 nM E.DNA + 9 (red) and E.DNA + 21 (blue) in the presence of 100 μM dNTP. All traces were fitted by Equation (1). Fitting of the gray trace according to Equation (1) provides $k_{obs1} = 19 \pm 1 \times 10^{-3}$ s⁻¹, $k_{obs2} = 17 \pm 3 \times 10^{-4}$ s⁻¹ and $a = 0.35 \pm 0.1$. Fitting of the red trace provides $k_{obs1} = 14 \pm 1 \times 10^{-3}$ s⁻¹, $k_{obs2} = 13 \pm 3 \times 10^{-4}$ s⁻¹ and $a = 0.85 \pm 0.1$ and fitting of the blue trace provides $k_{obs1} = 18 \pm 2 \times 10^{-2}$ s⁻¹, $k_{obs2} = 63 \pm 5 \times 10^{-4}$ s⁻¹ and $a = 0.5 \pm 0.1$. Excitation and emission wavelengths were 540 and 580 nm for the kinetic traces, respectively.

Figure 1. Design and validation of the FRET based assay. (A) Schematic representation of the FRET-based assay. The DNA–DNA p/t was labeled with both TMR (green) and Cy5 (red). Addition of RT and dNTP led to extension of the DNA primer, which shifted the two dyes further apart from each other, resulting in a decrease in FRET. (B) Polymerization kinetic trace of 100 nM RT with 100 nM doubly-labeled p/t duplex on addition of 100 μM dNTPs (red trace). The black line corresponds to the fit of the trace by Equation (1), with $k_{obs1} = 19 \pm 1 \times 10^{-3}$ s⁻¹, $k_{obs2} = 17 \pm 3 \times 10^{-4}$ s⁻¹ and $a = 0.35 \pm 0.1$. (Inset) Emission spectra of the doubly-labeled duplex in the absence (black line) and the presence (red line) of RT and after completion of polymerization with the addition of 100 μM dNTPs (blue line). For comparison, the emission spectrum of the duplex with TMR-labeled template and non-labeled primer (pink line) in the presence of RT is given to indicate the emission of the donor alone. Excitation and emission wavelengths for the kinetic trace were 540 and 580 nm, respectively.

Figure 2. Kinetic curves with primers of different lengths. (A) Nucleic acid substrate sequences used in this study. Acceptor and donor fluorophores are attached by a linker to the T residues represented in red and green, in the primer and template sequences, respectively. (B) Polymerization kinetic trace of 100 nM RT labeled in its p51 thumb subdomain by Alexa 488 with 100 nM duplex, where the Cy5-labeled primer is annealed to the template labeled at its 3' end by TMR, upon addition of 100 μM dNTPs. The black line corresponds to the fit of the trace by Equation (1), with $k_{obs1} = 17 \pm 3 \times 10^{-3}$ s⁻¹, $k_{obs2} = 9 \pm 2 \times 10^{-3}$ s⁻¹ and $a = 0.5 \pm 0.1$. Excitation and emission wavelengths were 480 and 520 nm, respectively. (C) Polymerization kinetic trace of 100 nM E.DNA (gray), 100 nM E.DNA + 9 (red) and E.DNA + 21 (blue) in the presence of 100 μM dNTP. All traces were fitted by Equation (1). Fitting of the gray trace according to Equation (1) provides $k_{obs1} = 19 \pm 1 \times 10^{-3}$ s⁻¹, $k_{obs2} = 17 \pm 3 \times 10^{-4}$ s⁻¹ and $a = 0.35 \pm 0.1$. Fitting of the red trace provides $k_{obs1} = 14 \pm 1 \times 10^{-3}$ s⁻¹, $k_{obs2} = 13 \pm 3 \times 10^{-4}$ s⁻¹ and $a = 0.85 \pm 0.1$ and fitting of the blue trace provides $k_{obs1} = 18 \pm 2 \times 10^{-2}$ s⁻¹, $k_{obs2} = 63 \pm 5 \times 10^{-4}$ s⁻¹ and $a = 0.5 \pm 0.1$. Excitation and emission wavelengths were 540 and 580 nm for the kinetic traces, respectively.

$\pm$
rates (46,48,53), indicating that the Cy5 label in the primer does marginally affect RT polymerization. An alternative hypothesis may be that the TMR-containing region in the template is progressively folded during primer extension (Figure 1A), giving complex changes in the distance and thus, in the FRET efficiency between TMR and Cy5, so that the observed fluorescence signal is no more directly related to the number of nucleotides incorporated. To check this hypothesis, we compared the elongation kinetic traces of E.DNA with those obtained with E.DNA + 9 and E.DNA + 21 primers (Figure 2A) that were extended by respectively 9 and 21 nt, in respect with E.DNA (Figure 2C). Interestingly, though the primer was only extended by 9 nt in E.DNA + 9, the initial donor fluorescence was already close that of the final plateau, which as expected, strongly affected the amplitude associated to $k_{obs1}$ (from 0.35 to 0.85). Moreover, the initial fluorescence of the trace with E.DNA + 21 was even higher than the plateau, indicating that the distance between the two dyes was increased as compared to the fully extended ds-DNA duplex. This increased distance likely results from the about 2-fold increased length per base of single stranded DNA as compared to double stranded DNA (56), so that at least for short segments, higher interchromophore distances can be achieved. Both the E.DNA + 9 and E.DNA + 21 data clearly indicate that the fluorescence changes observed as RT elongates the p/t DNA results from complex donor–acceptor distance changes, so that the apparent rate constants retrieved from the experimental curves do not reflect the real catalytic rates of RT.

Next, we explored the dependence of the $k_{obs}$ values on the dNTP concentrations to further identify the mechanistic steps monitored by this assay (Supplementary Figure S4). From the analysis of this dependence, the assay is thought to monitor the formation of the final extended DNA (56), so that at least for short segments, higher inter-chromophore distances can be achieved. Both the E.DNA + 9 and E.DNA + 21 data clearly indicate that the fluorescence changes observed as RT elongates the p/t DNA results from complex donor–acceptor distance changes, so that the apparent rate constants retrieved from the experimental curves do not reflect the real catalytic rates of RT.

Use of the assay to monitor the effect of RT inhibitors

As the assay monitors the RT polymerization, we next evaluated its ability to screen for RT inhibitors. To this end, we monitored the effect of two well-characterized RT inhibitors, Nevirapine and Zidovudine (AZTTP) (5–7,57), on RT polymerization in the presence of RT inhibitors. Nevirapine calculated from E.DNA complex (Figure 3C and D). $IC_{50}$ values determined for AZTTP were found to be 7 (±1) µM and 6 (±1) µM for $k_{obs1}$ and $k_{obs2}$, respectively. These values are in good agreement with the values reported in literature (48,59).

Similar dose-dependent decrease in the values of $k_{obs1}$ and $k_{obs2}$ was observed upon addition of Nevirapine to E.DNA complex (Figure 3C and D). $IC_{50}$ values for Nevirapine calculated from $k_{obs1}$ (Figure 3D) and $k_{obs2}$ (Figure 3D, inset) were found to be 71 (±8) nM and 74 (±9) nM, respectively. Once again, these $IC_{50}$ values were similar to previously reported $IC_{50}$ values (48,60). Thus, the assay appears highly relevant for quantitatively determining the potency of RT inhibitors.

Finally, in order to evaluate the quality of the assay for high-throughput screening (HTS) applications, we calculated a Z-factor for the $k_{obs1}$ and $k_{obs2}$ values at each inhibitor concentration, using Equation (3):

$$Z = 1 - 3 \times \frac{SD_{Sample}}{mean_{Sample} - mean_{Control}}$$

$$+ 3 \times \frac{SD_{Control}}{mean_{Sample} - mean_{Control}}$$

(Figure 3B) as a function of the inhibitor concentrations, $I$, were then used to determine $IC_{50}$ values, according to (58):
Here, we describe an efficient, robust and reproducible real-time assay to screen for RT inhibitors, by monitoring RT polymerization through changes in the FRET signal of a doubly-labeled p/t duplex. In contrast to most existing assays (28–42), this assay is a continuous one-step and one-pot method that does not require cumbersome sample handling. Moreover, it presents the advantage to use commercially available fluorescently labeled oligonucleotides and non-modified RT. This is in contrast to previous assays that require fluorescent or radiolabeled nucleosides (28–42) or chaperone activity of NCp7 (69). The addition of WDO-217 led to a full inhibition of the NCp7 effect on RT polymerization, providing the same kinetic parameters and fluorescence plateau as compared to RT alone (Figure 5A, compare magenta and green traces, and Figure 5B). In contrast, WDO-217 showed no effect on RT (Figure 5A, compare green and dark blue traces). The inhibition of NCp7 activity likely results from the previously demonstrated change in its binding mode to nucleic acids, as a result of zinc ejection (69). The binding mode of the zinc-depleted NCp7 is likely dominated by electrostatic interactions with the phosphate groups of the oligonucleotide backbone and is thought to prevent the nucleic acid chaperone properties of NCp7 as well as its productive interaction with RT. This has been verified by replacing NCp7 with NCp7(SSHS), a mutant that is unable to bind zinc (70) but still binds nucleic acids with high affinity (71). This mutant provided similar polymerization profile (Figure 5A, black trace) and kinetic parameters (Figure 5B) as compared to RT alone, mimicking the condition where WDO-217 is added to NCp7. Unfortunately, the additional control using an inhibitor of both proteins could not be performed as no molecule to our knowledge has been reported to be active both on RT polymerization and NCp7.

Taken together, our data indicate that our direct and continuous assay can monitor the concerted activity of NCp7 and RT, and has the potential to identify molecules that inhibit one of these proteins or both at the same time.

CONCLUSION

Here, we describe an efficient, robust and reproducible real-time assay to screen for RT inhibitors, by monitoring RT polymerization through changes in the FRET signal of a doubly-labeled p/t duplex. In contrast to most existing assays (28–42), this assay is a continuous one-step and one-pot method that does not require cumbersome sample handling. Moreover, it presents the advantage to use commercially available fluorescently labeled oligonucleotides and non-modified RT. This is in contrast to previous assays that require fluorescent or radiolabeled nucleosides (28–42) or chaperone activity of NCp7 (69). The addition of WDO-217 led to a full inhibition of the NCp7 effect on RT polymerization, providing the same kinetic parameters and fluorescence plateau as compared to RT alone (Figure 5A, compare magenta and green traces, and Figure 5B). In contrast, WDO-217 showed no effect on RT (Figure 5A, compare green and dark blue traces). The inhibition of NCp7 activity likely results from the previously demonstrated change in its binding mode to nucleic acids, as a result of zinc ejection (69). The binding mode of the zinc-depleted NCp7 is likely dominated by electrostatic interactions with the phosphate groups of the oligonucleotide backbone and is thought to prevent the nucleic acid chaperone properties of NCp7 as well as its productive interaction with RT. This has been verified by replacing NCp7 with NCp7(SSHS), a mutant that is unable to bind zinc (70) but still binds nucleic acids with high affinity (71). This mutant provided similar polymerization profile (Figure 5A, black trace) and kinetic parameters (Figure 5B) as compared to RT alone, mimicking the condition where WDO-217 is added to NCp7. Unfortunately, the additional control using an inhibitor of both proteins could not be performed as no molecule to our knowledge has been reported to be active both on RT polymerization and NCp7.

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the preparation and purification of fluorescently-labeled RT (43–47). Using AZT and Nevirapine as representative examples of NRTI and NNRTI, respectively, it was found that both compounds dramatically decreased the kinetic rate constants and the fluorescence plateau of the FRET-based progress curves. From the dependence of the kinetic rate constants on the inhibitor concentration, reliable IC$_{50}$ values were obtained for both inhibitors. Moreover, the assay shows high Z-factor values, showing its high sensitivity and suitability for HTS. In addition, the assay proved useful to monitor the promotion by NCp7 of RT polymerization. This promotion was fully reverted by the addition of an NCp7 inhibitor, indicating that this assay can also disclose NCp7 inhibitors. Therefore, the assay appears highly potent, being able to screen for RT and NCp7 inhibitors, as well as for compounds targeting the two proteins. The latter would be of high interest, as RT inhibitors are prone to resistance while due to its high conservation, NCp7 inhibitors are thought to be much less prone to resistance (15,16).

Finally, due to its simplicity and ease to use, our one-pot and one-step FRET-based assay will certainly be useful for monitoring the activity of other eukaryotic and prokaryotic polymerases, in order to substitute previous assays using more sophisticated fluorescence-based techniques (72–77) or radiolabeled substrates (78–80). Of course, the assay is expected also to present some limitations, the main one being the interference by the auto-fluorescent compounds in the libraries used for screening. Nevertheless, as we are using TMR and CY5 that absorb above 500 nm, the number of such interfering compounds should be limited. Another limitation will be provided by DNA intercalators that will alter the RT polymerization activity, and thus appear as false positives.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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