Biochemical, inhibition and inhibitor resistance studies of xenotropic murine leukemia virus-related virus reverse transcriptase

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ABSTRACT

We report key mechanistic differences between the reverse transcriptases (RT) of human immunodeficiency virus type-1 (HIV-1) and of xenotropic murine leukemia virus-related virus (XMRV), a gammaretrovirus that can infect human cells. Steady and pre-steady state kinetics demonstrated that XMRV RT is significantly less efficient in DNA synthesis and in unblocking chain-terminated primers. Surface plasmon resonance experiments showed that the gammaretroviral enzyme has a remarkably higher dissociation rate ($k_{off}$) from DNA, which also results in lower processivity than HIV-1 RT. Transient kinetics of mismatch incorporation revealed that XMRV RT has higher fidelity than HIV-1 RT. We identified RNA aptamers that potently inhibit XMRV, but not HIV-1 RT. XMRV RT is highly susceptible to some nucleoside RT inhibitors, including Translocation Deficient RT inhibitors, but not to non-nucleoside RT inhibitors. We demonstrated that XMRV RT mutants K103R and Q190M, which are equivalent to HIV-1 mutants that are resistant to tenofovir (K65R) and AZT (Q151M), are also resistant to the respective drugs, suggesting that XMRV can acquire resistance to these compounds through the decreased incorporation mechanism reported in HIV-1.

INTRODUCTION

Xenotropic murine leukemia virus-related virus (XMRV) is a gammaretrovirus that was first identified in some prostate cancer tissues (1,2) While some subsequent reports confirmed the presence of XMRV in prostate cancer samples (3–6), several others found little or no evidence of the virus in patient samples (7–9). XMRV DNA was also reported in 67% of patients with chronic fatigue syndrome (CFS) (10), but several subsequent studies in Europe and the USA failed to identify XMRV DNA in CFS patients or healthy controls (11–15). Hence, the relevance of XMRV to human disease remains unclear (16) and have been challenged (17). Most recently, it has been reported that XMRV has been generated through recombination of two separate proviruses suggesting that the association of XMRV with human disease is due to contamination of human samples with virus originating from this recombination event (18). Nonetheless, as a retrovirus that can infect human cells, XMRV can be very helpful in advancing our understanding of the mechanisms of retroviral reverse transcription, inhibition and drug resistance.

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XMRV RT is similar to the Moloney murine leukemia virus (MoMLV) RT, which has been the subject of structural and biochemical studies (19–24). Most of the differences between these gammaretroviral enzymes are at the RNase H domain (Supplementary Figure S1). Comparisons of human immunodeficiency virus type-1 (HIV) RT with MoMLV RT have revealed structural and sequence differences (21). For example, HIV-1 RT is a heterodimer composed of two related subunits (25,26) [reviewed in (27,28)]. Its larger p66 subunit (~66 kDa) contains both the polymerase and RNase H domains; the smaller p51 subunit, (~51 kDa), is derived from the p66 subunit by proteolytic cleavage and its role is to provide structural support and optimize RT’s biochemical functions (29). In contrast, structural studies have demonstrated that MoMLV RT is a monomer of about 74 kDa, although one study reported that it may form a homodimer during DNA synthesis (30). So far, there are no published biochemical or structural studies on XMRV RT. Hence, the present study on this enzyme and its comparison to related enzymes provides an excellent opportunity to advance our biochemical understanding of the mechanism of reverse transcription, its inhibition and drug resistance.

MATERIALS AND METHODS

Expression and purification of XMRV, HIV-1 and MoMLV RTs

The plasmid pBSK-XMRV containing the coding sequence of XMRV RT from the VP62 clone (GenBank: DQ399707.1) was chemically synthesized and optimized for expression by Epoch Biolabs Inc (Missouri City, Texas, USA). The 2013 bp XMRV RT sequence was amplified from pBSK-XMRVRT by PCR, using the forward and reverse primers 2 and 3. The digested amplicons were ligated into pET-28a (Novagen), resulting in NdeI and HindIII restriction sites. Drug resistant XMRV RT mutants Q190M and K103R (equivalent to HIV-1 Q151M and K65R) were generated by site-directed mutagenesis using forward and reverse primers 2 and 3. The digested plasmids were transformed into E. coli (Invitrogen) grown at 37°C. A cell pellet from a 3 l culture was induced with 150 μM IPTG at OD600 0.8, followed by 10 minutes with increasing amounts of MoMLV or HIV-1 RT for XMRV and MoMLV RTs, respectively and varying concentrations for HIV-1 RT were carried out in Reaction Buffer with 6 mM MgCl2 or 1.5 mM MnCl2, 0.5 mM EDTA, 200 nM or 100 nM Td26/5-Cy3-Pd18a, 20 nM or 5 nM RT for XMRV and MoMLV RTs, respectively and varying concentrations of dNTP in a final volume of 10 μl. The reactions for HIV-1 RT were carried out in Reaction Buffer with 100 nM Td26/5-Cy3-Pd18b, 10 nM HIV-1 RT and 6 mM MgCl2 in a 20 μl reaction. All the concentrations mentioned here and in subsequent assays reflect final concentration of reactants otherwise mentioned reactions were stopped after 15 min for XMRV, 4 min for MoMLV RT, and 2.5 min for HIV-1. The products were resolved on native 6% polyacrylamide–5M urea gels. The gels were scanned with a Fuji Fla-5000 PhosphorImager (Stamford, CT, USA) and the bands were quantified using MultiGauge. Results were plotted using GraphPad Prism 4.

Steady state kinetics

Steady state parameters $K_m$ and $k_{cat}$ for dATP incorporation were determined using single nucleotide incorporation gel-based assays. XMRV RT and MoMLV RT reactions were carried out in 50 mM Tris–HCl pH 7.8, 60 mM KCl, 0.1 mM DTT, 0.01% bovine serum albumin (BSA) (Reaction Buffer) with 6 mM MgCl2 or 1.5 mM MnCl2, 0.5 mM EDTA, 200 nM or 100 nM Td26/5-Cy3-Pd18b, 20 nM or 5 nM RT for XMRV and MoMLV RTs, respectively and varying concentrations of dNTP in a final volume of 10 μl. The reactions for HIV-1 RT were carried out in Reaction Buffer with 100 nM Td26/5-Cy3-Pd18b, 10 nM HIV-1 RT and 6 mM MgCl2 in a 20 μl reaction. All the concentrations mentioned here and in subsequent assays reflect final concentration of reactants otherwise mentioned reactions were stopped after 15 min for XMRV, 4 min for MoMLV RT, and 2.5 min for HIV-1. The products were resolved on 15% polyacrylamide–7M urea gels. The gels were scanned with a Fuji Fla-5000 PhosphorImager and the bands were quantified using MultiGauge. Results were plotted using GraphPad Prism 4. $K_m$ and $k_{cat}$ were determined graphically using Michaelis–Menten equation.

Gel mobility shift assays

Formation of RT-DNA binary complex: 20 nM Td31/5-Cy3-Pd18a (Supplementary Table S1) was incubated for 10 minutes with increasing amounts of MoMLV or XMRV RT in 50 mM Tris–HCl pH 7.8, 0.01% BSA, 5 mM MgCl2 and 10% (v/v) sucrose. The complexes were resolved on native 6% polyacrylamide 50 mM Tris–borate gel and visualized as described above.

Active site titration and determination of $K_{D,DNA}$

Active site concentrations and kinetic constants of DNA binding for XMRV, HIV-1 and MoMLV RTs were determined using pre-steady state experiments. Reactions...
with XMRV and MoMLV RTs were carried out in the reaction buffers listed above. For XMRV RT 100nM protein was pre-incubated with increasing concentrations of Td31/5'-Cy3-Pd18a, followed by rapid mixing with a reaction mixture containing 5mM MgCl2 and 100μM next incoming nucleotide (dTTP). The reactions were quenched at various times (5ms to 4s) by adding EDTA to a final concentration of 50mM. The amounts of 19-mer product were quantified and plotted against time. The data were fit to the following burst equation:

\[ P = A(1 - e^{-kt}) + k_{ss}t \]  

where \( A \) is the amplitude of the burst phase that represents the RT–DNA complex at the start of the reaction, \( k_{obs} \) is the observed burst rate constant for dNTP incorporation, \( k_{ss} \) is the steady state rate constant and \( t \) is the reaction time. The rate constant of the linear phase (\( k_{cal} \)) was estimated by dividing the slope of the linear phase by the enzyme concentration. The active site concentration and T/P binding affinity (\( K_{D,RT} \)) were determined by plotting the amplitude (\( A \)) against the concentration of T/P. Data were fit to the quadratic equation (Equation 2) using non-linear regression:

\[ A = 0.5(K_D+|RT|+|DNA|) - \sqrt{0.25(K_D+|RT|+|DNA|)^2 - ((|RT|+|DNA|) - (|RT|+|DNA|))} \]  

where \( K_D \) is the dissociation constant for the RT–DNA complex, and [RT] is the concentration of active polymerase. HIV-1 RT’s DNA binding affinity was determined as previously described (29).

**Surface plasmon resonance assay**

We used surface plasmon resonance (SPR) to measure the binding constants of XMRV and HIV-1 RTs to double-stranded DNA. Experiments were carried out using a Biacore T100 (GE Healthcare). To prepare the sensor chip surface we used the 5'-biotin-Td31/Pd25 oligonucleotide (Supplementary Table S1). One hundred and twenty RUs of this DNA duplex were bound in channel 2 of a streptavidin-coated sensor chip [Series S Sensor Chip SA (certified)] by flowing a solution of 0.1μM DNA at a flow rate of 10μl/min in a buffer containing 50mM Tris pH 7.8, 50mM NaCl. The binding constants were determined as follows: RT binding was observed by flowing solutions containing increasing concentrations of the enzyme (0.2, 0.5, 1, 2, 5, 10, 20, 50, 100 and 200nM) in 50mM Tris pH 7.8, 60mM KCl, 1mM DTT, 0.01% NP40 and 10mM MgCl2 in channels 1 (background) and 2 (test sample) at 30μl/min. The trace obtained in channel 1 was subtracted from the trace in channel 2 to obtain the binding signal of RT. This signal was analyzed using the Biacore T100 Evaluation software to determine \( K_{D,RT} \), \( k_{on} \) and \( k_{off} \).

**Pre-steady state kinetics of dNTP incorporation**

The optimal nucleotide incorporation rates (\( k_{pol} \)) were obtained by pre-steady state kinetics analysis using single nucleotide incorporation assays. A solution containing XMRV RT (150nM final concentration) and Td31/5'-Cy3-Pd18a (40nM) was rapidly mixed with a solution of MgCl2 (5mM) and varying dATP (5–200μM) for 0.1 to 6s before quenching with EDTA (50mM) (all concentrations in parentheses are final, unless otherwise stated). Products were resolved and quantified as described above. Burst phase incorporation rates and substrate affinities were obtained from fitting the data to Equation 1. Turnover rates (\( k_{pol} \)), dNTP binding to the RT-DNA complex (\( K_{D,dNTP} \)), and observed burst rates (\( k_{obs} \)) were fit to the hyperbolic equation:

\[ k_{obs} = (k_{pol}[dNTP])/(K_{D,dNTP}+[dNTP]) \]  

HIV-1 RT’s DNA binding affinity was determined as previously described (29).

**Fidelity of DNA synthesis**

The fidelity (error-proneness) of XMRV RT was determined and compared with that of MoMLV RT and HIV-1 RT by primer extension assays using 10nM heteropolymeric Td100/5'-Cy3-Pd18a. Reactions (10μl) were carried out in Reaction Buffer containing all four dNTPs (100μM each) or only three dNTPs (missing either dATP, dGTP or dTTP) at 100μM each. Incubations of the XMRV and MoMLV (50nM) reactions were at 37°C for 45min and 30min for HIV-1 RT (20nM). Reactions were initiated by adding dNTPs, stopped with equal volume of formamide-bromophenol blue, and an aliquot was run on a 16% polyacrylamide–7M urea gel.

**Kinetics of mismatch incorporation**

For these experiments, instead of including the next correct nucleotide (dATP) in the polymerase reactions, we used dTTP as the mismatched incoming nucleotide. Hence, 50nM XMRV RT was pre-incubated with 35nM Td31/5'-Cy3-Pd18a in reaction mixture. Reactions were initiated by adding dTTP (5–750μM) and 5mM MgCl2 followed by incubation (37°C) for 5min, due to the decreased mismatch incorporation rate of XMRV. For MoMLV RT, 30nM RT and 20nM DNA used and the reactions were carried out for 2.5 minutes. For HIV-1, 30nM RT, 20nM DNA and 0–200μM nucleotide were used and the reactions were carried out for 2.5 min. The amount of extended primer was quantified and plotted against the concentration of dTTP. The data were used to derive the \( K_{D,dNTP} \) of incorrect nucleotide binding, the rate \( k_{pol} \) (using Equations 1 and 3) and the efficiency of the misincorporation reaction (\( k_{pol}/K_{D,dTTP} \)).

**Determination of in vivo fidelity**

ANGIE P cells, which contain a retroviral vector (GA-1) that encodes a bacterial \( \beta \)-galactosidase gene (lacZ) and a neomycin phosphotransferase gene, were plated (5 x 10⁶ cells/100 mm dish) and after 24h were transfected using the calcium phosphate precipitation method with a plasmid expressing either XMRV or amphotropic MLV (AM-MLV) (three independent transfections per vector). After 48h, the culture medium with XMRV or (AM-MLV) was harvested, serially diluted and used to infect
D17 target cells (2 x 10^5 cells/60 mm dish) in the presence of polybrene. The infected D17 cells were selected for resistance to G418 (400 μg/ml) in the presence of 1 μM AZT to suppress recombination, and characterized by staining with 5-bromo-4-chloro-3-indolyl-β-D-galacto-pyranoside (X-Gal) ~2 weeks after G418 selection. The frequencies of inactivating mutations in lacZ quantified as described before (blue versus white colonies) (34).

**Processivity of DNA synthesis—trap assay**

Processivity reactions were carried out in Reaction Buffer containing 20 nM T_d100/P_d18, 100 μM of each dNTP, 30 nM HIV-1 RT, 50 nM MoMLV RT or 100 nM XMRV RT and 1 μg/ml unlabeled calf thymus DNA trap in 50 μL. Enzymes were pre-incubated with T_d100/P_d18 for 1 min before adding dNTPs (100 μM each) together with the calf thymus DNA trap. Reactions were incubated at 37°C, and 10 μl aliquots were taken out at 3, 7.5 and 15 min for HIV-1 RT or at 7.5, 15 and 30 min for XMRV RT and MoMLV RT, and mixed with equal volume of loading dye. The effectiveness of the trap was determined by pre-incubating the enzyme with the trap before adding T_d100/P_d18. Control DNA synthesis was measured in absence of trap under the same conditions. Reaction products were resolved and quantified as described above. Bands corresponding to full extension products were quantified using MultiGauge Software and IC_{50}s were obtained from dose–response curves using GraphPad Prism.

**PPi- and ATP-dependent excision and rescue of T/P_{AZT-MP} or T/P_{EFdA-MP}**

The ability of enzymes to use PPi or ATP to unblock template-primers that had AZT-MP (T/P_{AZT-MP}) or EFdA-MP (T/P_{EFdA-MP}) at their 3’ primer ends was measured as follows: 20 nM of T/P_{AZT-MP} or T/P_{EFdA-MP} were prepared as described before (32). They were incubated at 37°C with either 60 nM HIV-1 RT or 200 nM XMRV RT in the presence of 0.15 mM PPi or 3.5 mM ATP for PPi- or ATP-dependent rescue reactions, respectively. Reactions were initiated by the addition of MgCl₂ (6 mM). Aliquots were removed at different times (0–90 min) and analyzed as above. Rescue assays were performed in the presence of 100 μM dATP to prevent EFdA-MP reincorporation, 0.5 μM dTTP, 10 μM ddGTP and 10 mM MgCl₂.

**Molecular modeling**

The sequence of XMRV RT from the VP62 clone was aligned with that of MoMLV RT (PDB: 1RW3) (21,22) using ClustalW. To generate the homology model of XMRV RT, we used the Prime protocol of the Schrödinger software suite (Schrödinger Inc. NY). The resulting molecular model was further energy minimized by OPLS2005 force field using the Impact option of Schrödinger. The final model was validated with PROCHECK v.3.5.4.

**RESULTS**

**Comparison of RT sequences**

The XMRV and MoMLV enzymes are closely related (~95% sequence identity) with most of the differences between them being in the RNase H domain (Supplementary Figure S1). While XMRV and MoMLV differ significantly from HIV-1 RT, the known polymerase motifs (A–F) are well conserved in all three enzymes (Supplementary Figure S1). Specifically, the active site aspartates in Motifs A and C (Figure 9) (D150, D224, D225 in XMRV RT; D150, D224, D225 in MoMLV RT; D110, D185, D186 in HIV-1 RT) are conserved in all three RTs. Also, the three enzymes are similar in Motif B, which is involved in dNTP binding and multidrug resistance (AZT and deoxynucleoside drugs) through the decreased incorporation mechanism (27,39–41). Specifically, all three enzymes have a glutamate at the start of this motif (Q151 in HIV-1 RT, Q190 in XMRV RT and Q190 in MoMLV RT). Motif D includes HIV-1 RT residues L210 and T215, which when mutated they enhance excision of AZT from template-primers (T/PAZT-MP or T/PEFdA-MP) that had AZT-MP (T/P_{AZT-MP}) or EFdA-MP (T/P_{EFdA-MP}) at their 3’ primer ends. These motifs are mostly conserved in HIV-1 RT, but differ from XMRV and MoMLV RTs, where the corresponding residues are N226 and A231 (Supplementary Figure S1). K219 of HIV-1 RT Motif D is proximal to resolving the products on 15% polyacrylamide–7 M urea gels and visualized as described above. Bands corresponding to full extension products were quantified using MultiGauge Software and IC_{50}s were obtained from dose–response curves using GraphPad Prism.

**Assays for reverse transcriptase inhibition**

DNA synthesis by 50 nM XMRV RT or MoMLV RT was carried out in Reaction Buffer using 20 nM T_d100/5’-Cy3-P_d18a, 2.5 μM dNTPs, 5 mM MgCl₂ and varying amounts of NRTI (0–100 μM). Reactions were quenched with 95% formamide after 1 h incubation at 37°C (38). In experiments with aptamers 10 nM XMRV RT, 20 nM T_d100/5’-Cy3-P_d18a and 50 μM dNTPs were used in the presence of varying amounts of aptamer for 30 min (0–500 nM for m.1.3; 0–25 nM for m.1.4 and m.1.1FL). The inhibition of DNA polymerization was monitored by resolving the products on 15% polyacrylamide–7 M urea gels and visualized as described above. Bands corresponding to full extension products were quantified using MultiGauge Software and IC_{50}s were obtained from dose–response curves using GraphPad Prism.
the dNTP-binding pocket and is also conserved in the other enzymes (K235). The DNA primer grip (Motif E) (36,42) in HIV-1 RT (M230G231,Y232) is slightly different in the gammaretroviral enzymes (L245G246,Y247). Motif F at the fingers subdomain of all enzymes has two conserved lysines that bind the triphosphate of the dNTP (K65 and K72 in HIV-1 RT; K103 and K110 in XMRV and MoMLV RTs).

Several HIV-1 residues involved in NRTI resistance have the resistance mutations in XMRV and MoMLV RTs (Table 1). Hence, XMRV and MoMLV RTs have a Val as the X residue (codon 223) of the conserved YXDD sequence of Motif C. An M184V mutation at this position in HIV-1 RT causes strong, steric hindrance-based, resistance to 3TC and FTC (43–45), and to a lesser extent to ddI, ABC [reviewed in (46)], and translocation defective RT inhibitors (TDRTIs) (43) (Table 1). Similarly, the M41L mutation, which causes excision-based AZT resistance in HIV is already present in XMRV and MoMLV RT (L81, Table 1). The gammaretroviral enzymes differ from HIV-1 RT in several other HIV drug resistance sites (HIV residues 62, 67, 69, 70, 75, 77, 115, 210, 215) (Table 1). Finally, there are also differences in residues that are essential for NNRTI binding in HIV-1 RT: W229 changes to Y268 in XMRV RT, Y181 to L220, Y188 to L227 and G190 to A229 (Table 1) (27,28,47–49).

Preparation of MoMLV and XMRV RTs

The sequence coding for full-length XMRV RT from the VP-62 clone (NCBI RefSeq: NC_007815) (1) was optimized for expression in bacteria, synthesized by Epoch Biolabs and cloned as described in ‘Materials and Methods’ section. Both XMRV RT and MoMLV RT were tagged with a hexahistidine sequence at the N-terminus and expressed with a yield of >2 mg/l of culture. Purified enzymes (>95% pure, Supplementary Figure S2) were stored at −20°C. The presence of NP-40 or glycerol was critical for enzyme stability.

Steady state kinetics of nucleotide incorporation

Initial polymerase activity assays using Td31/5'-Cy3-Pd18a displayed overall slower polymerase activity of XMRV RT compared to HIV-1 and MoMLV RTs. This observation led us to investigate the steady state nucleotide incorporation properties of XMRV RT using single nucleotide incorporation assays. The estimated values for $k_{cat}$ (19.9 min$^{-1}$ for HIV-1 RT (32), 3.3 min$^{-1}$ for MoMLV RT, 0.6 min$^{-1}$ for XMRV RT) and $K_m$ (mM for HIV-1 RT, 3.3 mM for MoMLV RT, 3.0 mM for XMRV RT) show that XMRV RT has a drastically reduced efficacy ($k_{cat}/K_m$) at nucleotide incorporation, compared to both MoMLV and HIV-1 RTs.

DNA binding affinity

To assess if the efficiency of XMRV RT was also affected by a lower DNA binding affinity we measured the DNA binding affinity of the enzymes using three methods: gel-mobility shift assays, pre-steady state kinetics and SPR. Gel-mobility shift assays showed that the $K_D$ for XMRV RT was marginally higher than that for HIV-1 RT and MoMLV RT (data not shown) (50) suggesting weaker binding to DNA.

DNA binding affinity using pre-steady state kinetics

Pre-steady state kinetics allows estimation of the fraction of active polymerase sites as well as the $K_D$ value for the enzyme. The amplitudes of DNA extensions using XMRV RT and/or MoMLV RT at varying DNA concentrations were plotted against the DNA concentration and

The HIV-1 RT data are based on data from the Stanford HIV Database (85). wt = wild-type.

### Table 1. HIV-1 RT drug resistance mutations with wild-type XMRV RT and MoMLV RT residues

| HIV-1 residue numbers | HIV-1 RT wt | HIV-1 resistance mutations | XMRV RT wt | MoMLV RT wt |
|-----------------------|-------------|----------------------------|------------|-------------|
| Thymidine analog mutations (TAMs) | 184 | M | V | V | V | V | V223 | V223 |
| | 41 | M | – | L | L | L | – | L81 | L81 |
| | 67 | D | – | N | N | N | – | G105 | G105 |
| | 210 | L | – | W | W | W | – | N226 | N226 |
| | 215 | T | – | FY | FY | FY | – | A231 | A231 |
| | 219 | K | – | – | – | – | – | K235 | K235 |
| Non-thymidine analog regimen mutations | 65 | K | RN | RN | RN | RN | – | K103 | K103 |
| | 70 | K | EG | EG | EG | EG | – | D108 | D108 |
| | 74 | L | – | VI | – | – | – | V112 | V112 |
| | 75 | V | – | TM | M | TM | – | Q113 | Q113 |
| | 115 | Y | – | F | F | F | – | F155 | F155 |
| Multi-NRTI resistance mutations | 69 | T | Ins | Ins | Ins | Ins | – | N107 | N107 |
| | 151 | Q | M | M | M | M | – | Q190 | Q190 |
| | 62 | A | V | V | V | V | – | P104 | P104 |
| | 75 | V | – | I | – | I | – | Q113 | Q113 |
| | 77 | F | – | L | – | L | – | L115 | L115 |
| | 116 | F | – | Y | – | Y | – | F156 | F156 |
| TDRTI Mutations | 184 | M | V | V | V | V | V223 | V223 |
| | 165 | T | – | – | – | R | H204 | H204 |
the data were fit to the quadratic equation (Equation 2), yielding a $K_{D,DNA}$ of 33 nM for XMRV RT, 19 nM for MoMLV RT (Table 2) and 12.5 nM for HIV-1 RT (32). These values did not change significantly when tested with DNA of different lengths (data not shown). Hence, the transient kinetic experiments confirmed the findings of the gel-mobility shift assays showing XMRV RT to have lower DNA binding affinity than HIV-1 RT.

### Binding kinetics of XMRV and HIV-1 RT to double-stranded DNA

Measurements of $K_{D,DNA}$ using gel-mobility shift assays and pre-steady state kinetic methods do not offer insights regarding the kinetics of binding and release of nucleic acid from the viral polymerases. Hence, we used SPR to measure directly DNA binding and the DNA dissociation constants of the $K_{D,DNA}$. We attached on the SPR chip a nucleic acid biotinylated at the 5’ template end and immobilized it on a streptavidin sensor chip. Various concentrations of either XMRV or HIV-1 RT were flowed over the chip to measure the association ($k_{on}$) and dissociation ($k_{off}$) rates of the enzymes in real time (Figure 1). HIV-1 RT had considerably slower dissociation rates than XMRV RT, and longer dissociation phases were needed to obtain reliable values.

Several methods were tested to best fit our data. The ‘heterogeneous ligand’ method gave the best fit for both XMRV and HIV-1 RT. In this model the $x^2$ values for DNA binding to XMRV and HIV-1 RT were 9.3 RU$^2$ and 48.1 RU$^2$, respectively, compared to 15.1 RU$^2$ and 152 RU$^2$ when we tried fitting the data in a ‘homogeneous ligand’ model. The former model assumes that RT binds DNA in two different modes and provides two association ($k_{on}$) and two dissociation constants ($k_{off}$).

Our data show that XMRV RT has a slightly faster rate of association ($k_{on}$) than HIV-1 RT. We measured two $k_{on}$ values of 7.3 x 10$^{-3}$ M$^{-1}$ s$^{-1}$ and 8.2 x 10$^{-3}$ M$^{-1}$ s$^{-1}$ for XMRV RT versus 7.6 x 10$^{-3}$ M$^{-1}$ s$^{-1}$ and 1.2 x 10$^{-3}$ M$^{-1}$ s$^{-1}$ for HIV-1 RT. Interestingly, the dissociation rate of XMRV RT was significantly faster than that of HIV-1 RT (0.28 s$^{-1}$ and 0.0045 s$^{-1}$ for XMRV RT and 7.8 x 10$^{-3}$ s$^{-1}$ and 0.0076 s$^{-1}$ for HIV-1 RT) (Table 3). This difference in dissociation rate resulted in a $K_{D,DNA}$ at least 1 order of magnitude higher for XMRV RT compared to HIV-1 RT (38 and 54 nM versus 1.0 and 6.1 nM for XMRV and HIV-1 RT, respectively) (Table 3).

### Table 2. Kinetic parameters of DNA binding and synthesis by HIV-1 and XMRV RTs

| Nucleotide affinity and incorporation | HIV-1 RT | MoMLV RT | XMRV RT |
|--------------------------------------|----------|----------|---------|
| $K_{d,dNTP}$ (µM)                   | 1.3 ± 0.4 | 25 ± 5.3 | 26.6 ± 6.5 |
| $k_{pol}$ (s$^{-1}$)                | 24.4 ± 0.9 | 14.1 ± 0.8 | 8.9 ± 0.6 |
| $k_{pol}/K_{d,dNTP}$ (s$^{-1}$·µM$^{-1}$) | 18.8 | 0.56 | 0.33 |
| DNA binding affinity: $K_{D,DNA}$ (nM) | 12.5 | 19.0 | 32.5 |

*aHIV-1 RT data published previously (29).*

### Nucleotide binding affinity and optimal incorporation efficiency

A transient-state kinetics approach was used to estimate the dNTP binding affinity ($K_{d,dNTP}$) and maximum nucleotide incorporation rate ($k_{pol}$) (55). The rates at varying concentrations of next incoming nucleotide (dATP) were determined by plotting the amount of extended primer as a function of time. The rates were then plotted against dATP concentration. The data were fit to a hyperbola (Equation 3). The $K_{d,dNTP}$ for XMRV RT is 26.6 µM and the $k_{pol}$ is 8.9 s$^{-1}$ (Figure 2) (Table 2). Under similar conditions the $K_{d,dNTP}$ and $k_{pol}$ were 1.3 µM and 24.4 s$^{-1}$ for HIV-1 RT (29) and 25 µM and 14.1 s$^{-1}$ for MoMLV RT.

![Figure 1. Assessment of $K_{D,DNA}$, $k_{on}$ and $k_{off}$ using surface plasmon resonance. SPR was used to measure the binding affinity of RTs to a nucleic acid substrate. Increasing concentrations of each RT (0.2, 0.5, 1, 2, 5, 10, 20, 50, 100 and 200 nM) were injected over a streptavidin chip with biotinylated double-stranded DNA immobilized on its surface as described in ‘Materials and Methods’ section. The experimental trace (red) shown is the result of a subtraction of the data obtained from the channel containing the immobilized nucleic acid minus the signal obtained from an empty channel. The black curve represents the fitted data according to the ‘heterogeneous ligand’ model that assumes two different binding modes for RT on the nucleic acid.](Image 336x332 to 553x533)

### Table 3. DNA binding constants for HIV-1 and XMRV RTs from surface plasmon resonance

|                        | HIV-1 RT | XMRV RT |
|------------------------|----------|---------|
| $k_{pol}$ (M$^{-1}$ s$^{-1}$) | 7.6 x 10$^3$ | 7.3 x 10$^6$ |
| $k_{off}$ (s$^{-1}$)     | 7.8 x 10$^{-4}$ | 2.8 x 10$^{-1}$ |
| $K_{D,DNA}$ (nM)        | 1        | 38 (38-fold)* |
| $k_{on}$ (M$^{-1}$ s$^{-1}$) | 1.2 x 10$^6$ | 8.2 x 10$^4$ |
| $k_{off}$ (s$^{-1}$)     | 7.6 x 10$^{-3}$ | 4.5 x 10$^{-3}$ |
| $K_{D,DNA}$ (nM)        | 6.1      | 54 (9-fold)* |

*Increase in $K_{D,DNA}$ (decrease in affinity) with respect to HIV-1 RT. ($K_{D1-XMRV RT}/K_{D1-HIV-1-RT}$ and $K_{D2-XMRV RT}/K_{D2-HIV-1-RT}$).
Fidelity of nucleotide incorporation

To assess whether XMRV RT displays high nucleotide incorporation fidelity we monitored the incorporation of three dNTPs by XMRV RT and compared with HIV-1 RT (52). The results of fidelity assay are shown in Figure 3. The lanes marked ‘4dNTPs’ for all enzymes represent the DNA synthesis using a Td100/5'-Cy3-Pd18a template-primer in the presence of all four dNTPs. The subsequent lanes, marked ‘-dNTP’, correspond to the synthesis of DNA in the absence of that specific deoxy-nucleotide triphosphate. The comparison of the DNA synthesis in the absence of one nucleotide by HIV-1 RT, MoMLV RT and XMRV RT shows that HIV-1 and MoMLV RTs were able to misincorporate and extend the primer beyond the missing nucleotide more efficiently than XMRV RT, suggesting that the latter is a less error prone DNA polymerase. It should be noted that the higher fidelity of XMRV is not the result of measuring a smaller number of errors because of the decreased replication rate, as the assay conditions were optimized to allow production of the same amount of full length product in the presence of all four dNTPs for and MoMLV RTs. To further investigate the fidelity of DNA synthesis by XMRV RT, the kinetics of mismatch nucleotide incorporation were carried out in a quantitative manner by monitoring the incorporation of single mismatched nucleotide under pre-steady state conditions. The estimated $K_{d,dNTP}$ (mismatch) and $k_{pol}$ values show that XMRV RT has a lower affinity for a mismatched nucleotide but comparable turnover number than MoMLV RT, suggesting that the observed higher fidelity over MoMLV RT is due to differences during the nucleotide-binding step (Table 4). However, compared to HIV-1 RT, XMRV RT has decreased both affinity and incorporation rate, suggesting that its higher fidelity is the result of both decreased binding of mismatched nucleotides and slow rate of incorporation.

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**Figure 2.** Pre-steady state kinetics of nucleotide incorporation by XMRV RT. 150 nM XMRV RT was pre-incubated with 40 nM Td100/5'-Cy3-Pd18a rapidly mixed with a solution containing MgCl₂ (5 mM) and varying concentrations of dATP: 25 μM (filled square), 35 μM (filled triangle), 50 μM (filled inverted triangle), 75 μM (filled rhombus), 100 μM (filled circle), 125 μM (open square) and 150 μM (open triangle); and incubated for 0.1 to 6 s before being quenched with EDTA. The DNA product for each dATP concentration was fit to the burst equation (A). The burst amplitudes generated for each dATP concentration were then fit to a hyperbola equation (B) yielding the optimal rates of dNTP incorporation; $k_{pol}$ (8.9 s⁻¹) and $dNTP$ binding to the RT-DNA complex; $K_{d,dNTP}$ (26.6 μM).

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**Figure 3.** Comparison of in vitro fidelity of HIV-1, MoMLV and XMRV RTs. Extension of 10 nM Td100/5'-Cy3-Pd18a by HIV-1 RT, MoMLV RT or XMRV RT (20, 50 and 50 nM, respectively) in the presence of 150 μM each of three out of four nucleotides (the missing nucleotide is marked at the bottom of each lane). Reactions were run for 30 min for HIV-1 RT and 45 min for XMRV RT and MoMLV RT. For each enzyme the first lane in each set shows the position of unextended primer, the second lane shows full extension in the presence of all four dNTPs, and each consecutive lane shows extension in the presence of three dNTPs. The arrows on the right mark the expected pauses based on the indicated composition of the template strand.

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**Table 4.** Kinetics of mismatch incorporation for HIV-1, MoMLV and XMRV RTs

| Enzyme | HIV-1 RT | MoMLV RT | XMRV RT |
|--------|----------|----------|---------|
| $K_{d,dNTP}$ (μM) | 9 ± 0.3 | 38.9 ± 11.6 | 256 ± 72 |
| $k_{pol}$ (s⁻¹) | 6.81 ± 1.2 | 0.16 ± 0.01 | 0.15 ± 0.018 |
| $k_{pol}/K_{d,dNTP}$ (s⁻¹·μM) | 0.756 | 0.0041 | 0.00058 |
| Fidelity $^{a}$ | 0.04 | 0.007 | 0.002 |

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$a$Fidelity is the ratio of the incorporation efficiency ($k_{pol}/K_{d,dNTP}$) of the mismatched nucleotide (dTTP) over that of the correct (dTTP) ($k_{pol}/K_{d,dNTP}$).
Intracellular fidelity by measuring LacZ mutation frequency

The ANGIE P cells used for this assay are a D17-based encapsidating cell line and contain an MLV-based retroviral vector (GA-1), which encodes a bacterial β-galactosidase gene (*lacZ*) and a neomycin phosphotransferase gene (*neo*). Replication fidelity is a measure of the frequency of *lacZ* inactivation and was determined by measuring *lacZ* non-expressing white colonies. The results show that the number of white colonies was not statistically different in the case of XMRV as compared to AM-MLV, suggesting that under these conditions the fidelity of XMRV is not significantly different than that of AM-MLV (Figure 4).

Processivity of DNA synthesis

Processivity is the probability of translocation of a polymerase along a template and predicts the number of cycles of nucleotide incorporation during one productive enzyme–DNA binding event. We assessed XMRV RT's processivity of DNA synthesis in comparison to HIV and MoMLV RTs using both a gel-based trap assay and a quantitative pre-steady state assay. In the gel-based assay, the enzymes were pre-incubated with template-primer, then the reaction was initiated by the addition of all four nucleotides together with calf thymus DNA, which was used as a trap to bind free enzyme dissociated from the substrate during the course of the reaction (38). The length of the DNA product is an inverse measure of termination probability, as previously described. As a control, we used lanes where no trap was present; establishing that the same amount of total polymerase activity (processive and non-processive) is provided in all cases. The results indicate that XMRV RT is less processive than HIV-1 and MoMLV RTs with shorter DNA product after 30 min of reaction in the presence of trap (Figure 5).

To measure processivity quantitatively we applied a single turnover processivity assay developed by Patel et al. (35) (Figure 6). In this assay, the rates of consecutive nucleotide incorporations under single turnover conditions are monitored. The rate of elongation incorporation (*k*₁) and the rate of processive DNA synthesis (*k*₂) (Equation 4) were calculated at several template positions for each enzyme. The ratio of the rate of processive DNA synthesis to the rate of nucleotide incorporation (*k*₂/*k*₁) is referred to as the processivity index (35). The absolute values of these constants for HIV-1 RT, XMRV and MoMLV RT and their ratios are collected in Table 5. XMRV RT is clearly the least processive for each extension product. The difference in processivity varies significantly depending on sequence or sequence context (decrease in processivity from 3-fold up to 10-fold). While the current data do not allow generalization of rules for pausing at specific sites, this clearly shows consistently that XMRV is not as efficient as MoMLV RT in polymerizing processively through ‘difficult spots’.

Susceptibility of XMRV RT to NRTIs, TDRTIs and NNRTIs

Previous studies have shown that XMRV is inhibited by some antivirals (53–56). However, the susceptibility of XMRV RT has not been tested against a wide variety of...
nucleoside RT inhibitors (NRTIs) that block replication by chain-terminating the primer, or by preventing translocation after their incorporation into the nascent DNA chain (TDRTIs) (32,57,58). In addition, the susceptibility of XMRV RT to non-nucleoside RT inhibitors (NNRTIs) or RNA aptamers that can be selected to block reverse transcriptases (59–63) has not been established.

Hence, we performed gel-based primer extension assays in the presence of various inhibitors. As shown in Table 6, most of the HIV-1 RT inhibitors also block XMRV RT with significantly varying IC₅₀s. The most potent inhibitors tested were ENdA (4’-ethynyl-2-amino-2’-deoxyadenosine) followed by EFdA. EFdA was also potent at inhibiting wild-type XMRV replication in cell culture with an EC₅₀ of 40 nM from three independent experiments (standard error was 10 nM).

Unlike HIV-1 RT, XMRV RT and MoMLV RT lack the two tyrosine residues (Y181 and Y188 in HIV-1 RT) (Supplementary Figure S1) that are known to contribute to NNRTI binding. Hence, the gammaretroviral enzymes were not inhibited by the NNRTIs tested (TMC-125 and efavirenz) (Supplementary Figure S3).

Susceptibility of XMRV RT to RNA aptamers

We also tested XMRV RT’s susceptibility to three independent RNA aptamers that had been previously selected against MoMLV RT (60). The aptamers inhibited XMRV RT to varying extents with IC₅₀s ranging from 2 to 52 nM (Figure 7). Most notable was the m.1.1FL aptamer which gave IC₅₀s of 2 and 4 nM for XMRV RT (Figure 7) and MoMLV RT respectively, without inhibiting HIV-1 RT (data not shown). These inhibition assays utilized truncated forms of aptamers m.1.3 and m.1.4 lacking the 5’ and 3’ segments, demonstrating that these segments are not required.

PPi-mediated excision activity of XMRV RT

A key mechanism of NRTI resistance in HIV-1 RT is based on inhibitor excision from the primer end, using a pyrophospholytic reaction (64,65). The pyrophosphate donor in vivo is likely to be ATP, although PPi can efficiently unblock NRTI-terminated primers. This excision activity is present in wild-type HIV-1 RT, and is enhanced in the presence of AZT-resistance mutations. We measured the ability of wild-type XMRV to unblock primers terminated with AZT or EFdA in the presence of PPi. We found that unlike HIV-1 RT that excised AZT-MP efficiently under these conditions, XMRV RT had considerably lower excision activity (Figure 8). Similar excision experiments where ATP was used instead of PPi showed that XMRV is very inefficient in ATP-based excision as compared to HIV-1 RT (data not shown).
Susceptibility of mutant XMRV RTs to AZT-TP and tenofovir-DP

The HIV-1 RT mutation Q151M confers resistance to AZT by enhancing discrimination of the nucleotide analog leading to its reduced incorporation (37,66–68). Another HIV-1 RT mutation, K65R, decreases susceptibility to tenofovir (69,70). Since AZT and tenofovir are potent inhibitors of XMRV (Table 6) (54–56), we wanted to investigate whether the XMRV RT mutant equivalents of HIV Q151M and K65R (XMRV Q190M and K103R) would confer XMRV RT resistance to AZT and tenofovir. We constructed these mutant clones and tested their susceptibility to AZT and tenofovir in the same manner as wild-type XMRV RT. Interestingly, Q190M XMRV RT has a decreased susceptibility to AZT (approximately 5-fold increase in the IC50). Similarly, the K103R XMRV RT mutant enzyme was less susceptible to tenofovir, increasing the IC50 by at least 2-fold.

Molecular model of XMRV RT

Given the significant sequence similarity between XMRV and MoMLV RTs, the resulting homology model of XMRV RT is highly similar to MoMLV RT (>1.5 Å rms) and of excellent quality. Since the input structure of MoMLV RT did not contain the RNase H domain of the enzyme, the XMRV RT model is also missing this domain. The molecular model of the polymerase domain of XMRV RT is shown in Figure 9. An alignment of the MoMLV RT crystal structure (22) with the XMRV RT homology model highlights the few changes in the polymerase domain of XMRV RT. These are L29 (P in MoMLV), Q234 (L in MoMLV), R238 (Q in MoMLV) and N422 (D in MoMLV). From these, residue 422 is located in the nucleic acid binding cleft and may contribute to differences in the interactions with nucleic acid substrate. However, most of the differences between the gammaretroviral enzymes are in their RNase H domains and also in the first 30 N-terminal residues of the polymerase domain, for which we do not have structural data.

Figure 7. Inhibition of XMRV RT by RNA aptamers. 10 nM XMRV RT was incubated with increasing amounts of RNA aptamer in Reaction Buffer for 5 min at 37°C followed by addition of 20 nM Td31/Cy3-Pd18a and 50 μM of each dNTP. (A) The reactions were stopped after 30 min and resolved on a polyacrylamide gel. The predicted secondary structures of each aptamer were generated by mfold. (B) The percent full extension was quantified for m.1.1FL (filled inverted triangle), m.1.3 (filled circle) and m.1.4 (filled square) and data points fit to one-site competition non-linear regression using GraphPad Prism 4 to calculate IC50. HIV-1 RT was not susceptible to m.1.1FL (open triangle). (Errors represent data deviation from the fit).

Figure 8. PPI-mediated unblocking of AZT-(A) and EfdA-(B) terminated DNA. About 20 nM of (A) AZT- or (B) EfdA-terminated Td31/Cy3-Pd18c (T/PAZT-MP or T/PEfdA-MP) was incubated with HIV-1 RT (60 nM) or XMRV RT (200 nM) in the presence of 150 μM PPI and 6 mM MgCl2. Aliquots of the reactions were stopped at different time points (0–90 min) and resolved on a 15% polyacrylamide-7M urea gel as described in the ‘Materials and Methods’ section.

Susceptibility of mutant XMRV RTs to AZT-TP and tenofovir-DP

The HIV-1 RT mutation Q151M confers resistance to AZT by enhancing discrimination of the nucleotide analog leading to its reduced incorporation (37,66–68). Another HIV-1 RT mutation, K65R, decreases susceptibility to tenofovir (69,70). Since AZT and tenofovir are potent inhibitors of XMRV (Table 6) (54–56), we wanted to investigate whether the XMRV RT mutant equivalents of HIV Q151M and K65R (XMRV Q190M and K103R) would confer XMRV RT resistance to AZT and tenofovir. We constructed these mutant clones and tested their susceptibility to AZT and tenofovir in the same manner as wild-type XMRV RT. Interestingly, Q190M XMRV RT has a decreased susceptibility to AZT (approximately 5-fold increase in the IC50). Similarly, the K103R XMRV RT mutant enzyme was less susceptible to tenofovir, increasing the IC50 by at least 2-fold.

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| Aptamer    | IC50 (nM) |
|------------|-----------|
| m.1.1FL    | 2 ± 0.15  |
| m.1.3      | 52 ± 0.27 |
| m.1.4      | 7 ± 0.16  |

Figure 7. Inhibition of XMRV RT by RNA aptamers. 10 nM XMRV RT was incubated with increasing amounts of RNA aptamer in Reaction Buffer for 5 min at 37°C followed by addition of 20 nM Td31/Cy3-Pd18a and 50 μM of each dNTP. (A) The reactions were stopped after 30 min and resolved on a polyacrylamide gel. The predicted secondary structures of each aptamer were generated by mfold. (B) The percent full extension was quantified for m.1.1FL (filled inverted triangle), m.1.3 (filled circle) and m.1.4 (filled square) and data points fit to one-site competition non-linear regression using GraphPad Prism 4 to calculate IC50. HIV-1 RT was not susceptible to m.1.1FL (open triangle). (Errors represent data deviation from the fit).

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| Aptamer    | IC50 (nM) |
|------------|-----------|
| m.1.1FL    | 2 ± 0.15  |
| m.1.3      | 52 ± 0.27 |
| m.1.4      | 7 ± 0.16  |
information since they were not included in the original crystal structure of MoMLV RT. The differences between XMRV RT and HIV-1 RT are very significant. Unlike the HIV enzyme, XMRV RT appears to be a monomer in solution. Moreover, alignment of the HIV-1 RT–DNA complex with XMRV RT based on their active sites at the palm subdomains shows that the thumb subdomain of XMRV RT would have to be repositioned to be able to accommodate nucleic acid.

DISCUSSION

Early studies reported the presence of XMRV in stromal cells from prostate cancer patient samples and also in CFS clinical samples. Some of the subsequent studies confirmed these findings whereas several others failed to identify XMRV in prostate cancer or in CFS patients, even when same samples were used (71). It was recently reported that human sample contamination with mouse DNA can occur frequently (17,72–74). Moreover, two coauthors from this study have recently demonstrated that XMRV is the product of recombination events between two MLV proviruses, suggesting that XMRV may not be relevant to human disease (18). Nonetheless, XMRV is still an important human retrovirus and comparisons with HIV can provide valuable insights into the fundamental mechanisms of DNA polymerization, RT inhibition and drug resistance, (75).

There is high degree of sequence similarity between the XMRV and MoMLV RTs (95% amino acid identity), and much less so with HIV-1 RT. Based on gel filtration experiments we conclude that unlike HIV-1 RT, but similar to MoMLV RT, XMRV RT exists in solution primarily as a monomer. We also included comparisons with HIV-1 RT in this study as it has been extensively studied and provides an excellent frame of reference.

We report here that there are significant differences in the DNA polymerization efficiency of the three enzymes. Although the polymerase active sites of the XMRV and MoMLV enzymes are almost identical, there is a considerable decrease in the efficiency of nucleotide incorporation by XMRV RT. Most differences in sequence are at the RNase H domain and are likely to affect polymerization by changing the positioning of DNA at the polymerase active site.

We have recently solved the crystal structure of the XMRV RNase H at high resolution (1.5Å) (pdb 3P1G) (Kirby, K.A. et al., submitted for publication). We observed major differences in affinity for nucleic acid that we determined with gel-mobility shift assays and with pre-steady state kinetics. SPR experiments dissected in more detail the specific defect of XMRV RT in binding DNA. Surprisingly, XMRV RT can associate very rapidly with DNA, even more so than HIV-1 RT (Figure 1 and Table 3). However, it dissociates from DNA much faster than the HIV enzyme, resulting in an overall reduced binding affinity. A possible reason for the fast association and dissociation rates of XMRV RT may be the apparent monomeric state, which might offer facile access to the nucleic acid binding cleft, although with less contacts and lower affinity than HIV-1 RT, which is a heterodimer (76,77). This high rate of XMRV RT dissociation from DNA likely contributes to the decreased processivity observed in our study, and may have consequences in the recombination rates of this virus.

Previous sequences of XMRV from prostate cancer tumors showed low variability, suggesting that the virus may have a high fidelity of replication (1,10). Our study demonstrated that HIV-1 RT and MoMLV RT incorporated mismatched nucleotides and extended past the mismatches more efficiently than XMRV RT. Pre-steady state kinetics established that the higher overall fidelity of XMRV RT over MoMLV RT is due to a lower affinity for mismatched nucleotides. When compared to HIV-1 RT, however, XMRV RT has differs in both the nucleotide binding and incorporation steps. Nonetheless, XMRV did not have higher fidelity than a related amphotropic MLV virus or HIV-1 in a cell-based assay. It is possible that the high dNTP concentration in dividing cells can suppress mismatching events. We have previously shown (39) that as nucleotide concentrations vary in different cell lines, this can affect viral susceptibility to NRTIs, and possibly in this case also incorporation of mismatched nucleotides. Additional cell-based studies using multiple cell lines and a large panel of viruses should provide a better understanding of the relation between in vivo and in vitro fidelity.

Early studies have reported susceptibility of XMRV to some antiretrovirals that have been used in the treatment of HIV infection (53–56). In those studies the compounds were tested at the virus level. To better understand the interactions of inhibitors at their RT target level we tested here the ability of these and several more compounds to block the polymerase activity of XMRV RT. We found that two TDRTIs, EfDa-TP and EnDa-TP were very potent RT inhibitors (IC50s: 0.43µM and 0.14µM, respectively). Unlike other NRTIs, these compounds have a 3’ OH group and are known to efficiently inhibit HIV replication by blocking translocation (32,58,78).
Preliminary experiments demonstrated that they also block XMRV RT by the same mechanism (data not shown).

In HIV, moderate resistance to EFDa is conferred by the emergence of the M184V mutation at the conserved X position of the conserved YXDD motif of the polymerase active site. Interestingly, XMRV and MoMLV RTs already have a valine (V223) at this position. This difference is likely to contribute to the better potency of EFDa against HIV-1 RT than XMRV RT or MoMLV RT (57,58). It may also contribute to the decreased ability of XMRV RT to unblock chain-terminated primers, as was also reported for M184V HIV-1 RT (79) and to the enhanced fidelity reported here for XMRV RT, which is also reminiscent of the previously reported high fidelity of M184V HIV-1 RT (80,81). Nonetheless, despite the presence of a Val in the YMDD motif of XMRV RT, which is also reported for M184V HIV-1 RT (79) and to the enhanced fidelity reported here for XMRV RT, it was shown that introducing the primary Q151M mutation, followed by secondary mutations F77L, A62V, V75I and F116Y resulted in an enzyme with decreased susceptibility to AZT. Hence, it appears that these residues can confer AZT resistance to XMRV by reduced incorporation of nucleotide analogs, as is the case in HIV-2 (41). At this point we do not know if introduction of as yet unknown mutations could endow XMRV RT with the ability to unblock chain-terminated nucleic acids. The details of the molecular mechanism of XMRV resistance to tenofovir and AZT are under investigation.

In conclusion, our study provides detailed biochemical analysis of the mechanisms of polymerization, inhibition, fidelity, processivity and drug resistance of XMRV RT and how it compares with the closely related enzyme MoMLV RT and the more distantly related HIV-1 RT. The findings enhance our understanding of the basic mechanisms of reverse transcription.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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