Inhibition of the Replication of Clinical Drug-Resistant HIV-1 Strains by Small Molecule Integrase Inhibitors M522 and M532

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Abstract

Objective: Integrase (IN) is an enzyme essential for HIV-1 replication that has been a target of antiretroviral drug therapy. Since emerging HIV-1 variants have frequently become clinically resistant to antiretroviral agents, it is necessary to develop alternative IN inhibitors.

Methods: We tested IN inhibitors, M522 and M532, against clinically resistant HIV-1 strains to antiretroviral drugs (AZT, non-nucleoside reverse transcriptase inhibitors, IN drug raltegravir, protease inhibitors); wild-type and clinical isolate from HIV-infected patients. We performed disintegration studies to show the interaction of M522 and M532 with the catalytic core domain of HIV-1 IN and time-of-drug-addition experiments to determine the inhibition step of viral replication. We tested selection of HIV-1 with M522 and M532 to examine the emergence of new drug resistant virus. CD4+ cell count was calculated for several groups of cells infected with HIV, cells treated and non-treated with M522 and M532 to evaluate their protective effect.

Results: M522 and M532 inhibited the replication of HIV-1 strains (wild-type; drug-resistant; clinical isolate from infected patients; and laboratory strains) with high potency. These inhibitors interacted with the catalytic core domain of HIV-1 IN and blocked its activity, prevented viral integration. M522 and M532 interfered with the viral replication precisely at the integration step. HIV-1 virus did not develop resistance to M522 and M532 for 20 viral passages (160 days). These IN inhibitors protected the infected cells from cytopathic effects and the CD4+ cell counts of these cells treated with M522 and M532 were found to be identical to those of the uninfected cells.

Conclusion: M522 and M532 are potent against clinical isolate from HIV-infected patients, wild-type and clinical resistant strains especially the relevant raltegravir-resistant virus. Development of M522 and M532 as new mutation-insensitive drugs aiming for the protection of CD4+ T-cells during HIV infection in the clinical trials is in progress.

Keywords: HIV-1; Integrase inhibitors; Raltegravir; Drug resistance; CD4+ cell count

Introduction

Integrase (IN) is a clinically validated target for the treatment of human immunodeficiency virus infections [1]. Drug-induced mutations have been major obstacles in developing effective IN inhibitors for HIV-1 virus infection, which were reported with the diketo acid compounds [2]. The second most promising anti-HIV-1 IN developed was the raltegravir drug. In 2007, U.S. Food and Drug Administration (FDA) approved raltegravir (Isentress) as the first IN drug for treatment of HIV infected patients [3]. Raltegravir is a strand transfer inhibitor and HIV-1 strains resistant to this commercial drug were reported in 2008 [4,5]. Three primary mutations were responsible for raltegravir resistance. The two most frequent primary mutations were at positions Q148 (Q148K/R/H) and N155 (N155H); whereas, a third primary mutation at Y143 (Y143R/C/H) was less common [1,6,7]. A secondary mutation at position G140 (G140S) combined with the primary mutation Q148K/R/H, significantly enhanced drug resistance in the virus. The development of drug resistance has become a major cause of failure among antiretroviral therapies for treating HIV-1 infection effectively [4,5]. FDA next approved two more integrase strand transfer inhibitors (INSTIs), dolutegravir (Tivicay) and elvitegravir (Vitekta). Dolutegravir, was the first second-generation of INSTI, and was approved in August 2013 for the treatment of antiretroviral-naïve and antiretroviral-experienced patients [8]. Subsequently on September 24, 2014, FDA approved elvitegravir (Vitekta) as another INSTI used in combination with other antiretroviral drugs (HIV protease inhibitor) for the treatment of HIV-1 infection in treatment-experienced adults. Regrettfully, research showed a F121Y mutation in HIV-1 again for these integrase strand transfer inhibitors, dolutegravir, elvitegravir and raltegravir [9]. In addition to these catalytic site inhibitors, recent reports of several allosteric IN inhibitors, series of 2-(quinolin-3-yl) acetic acid derivatives (LEDGINs), established them as non-catalytic site inhibitors of HIV-1 IN. These compounds inhibited the interaction of LEDGF/p75-integrase. Studies again showed drug resistance in selection experiments with LEDGINs in cell culture [10-14]. Due to the failure of antiretroviral drugs and emergence of resistances to raltegravir, dolutegravir, elvitegravir, LEDGINs and other antiretroviral inhibitors, it is important to continue seeking IN inhibitors that are non-toxic to human cells and inhibit drug-resistant strains of HIV-1. Two such potent and selective HIV-1 IN inhibitors (M522 and M532) were previously isolated from the natural plant...
product, *Salvia miltiorrhiza* and they did not show cytoxicity at high concentrations [15].

HIV-1 IN is composed of three distinct domains, the N-terminal (residues 1–49), the catalytic core (residues 50–212), and the C-terminal (residues 213–288). The N-terminal domain is involved in enzyme multimerization, while the C-terminal domain, also called the DNA-binding domain, has strong but nonspecific DNA-binding activity [16]. HIV-1 IN, which mediates the integration of HIV-1 DNA and is required for stable production of provirus genes [17], has no mammalian counterpart, making it an attractive target for antiviral drug design [18]. HIV-1 IN catalyzes two well-characterized reactions, terminal 3′-end processing and strand transfer [16]. In addition, IN is capable of catalyzing the reverse reaction called disintegration [19]. This activity can be assayed in vitro to study the IN enzyme and its effects on viral integration, including the location of the enzyme’s catalytic core domain, and provides a convenient tool for measuring drug effects on IN deletion mutants (INΔ9–212) [19,20]. We applied this method in the present study to determine whether M522 and M32 interact with the catalytic core domain.

In the current study, we further characterized the antiviral activities of these two IN inhibitors against wild-type virus; different resistant strains of HIV-1 to reverse transcriptase (RT), protease (PR), and integrase (IN) drugs especially raltegravir-resistant strain as well as clinical isolate from HIV-infected patients. We performed disintegration experiments to determine the interaction of M522 and M32 with the catalytic core domain of HIV-1 IN. The protective values of M522 and M32 against HIV-1 were evaluated by comparing the CD4+ cell count in the drug treated infected cells and in the infected cells without treatments.

**Methods**

**Cells, viruses, and inhibitors**

We obtained H9, Sup-T1 cells and HIV-1 viruses (drug-resistant isolates and laboratory strains) from the AIDS Research and Reference Reagent Program, Division of AIDS, National Institute of Allergy and Infectious Diseases. In the present study, we used different drug-resistant isolates. HIV-1ΔRT/Gag, is an RT-resistant virus, HIV-1ΔINT/A17 variant, is highly resistant to inhibition by NNRTIs, and HIV-1ΔINT/M46SI/L53P/V287I/G214V is resistant to structurally diverse PR inhibitors. The two laboratory strains used were HIV-1ΔRT and HIV-1ΔINT. In addition, we used two other HIV-1 strains, a resistant one to the IN drug raltegravir harboring double mutations of G140S and Q148H and a wild-type strain (HIV-1NL4.3). Professor Zeger Debyser, Division of Molecular Medicine, Katholieke Universiteit Leuven, Flanders, Belgium generously provided these two strains. The drugs used were AZT (Retrovir), raltegravir (Isentress), and IN inhibitor (118-D-24), which were obtained from the Division of AIDS (NIAID, NIH). Dextran sulfate 5000 (DS), was purchased from Sigma-Aldrich Company and inhibitors M522 (Lithospermic acid) and M32 (lithospermic acid B) used in the current study were purified from *Salvia miltiorrhiza* roots as previously described [15].

**Viral infection and drug treatment**

H9 and Sup-T1 cells were grown and maintained in RPMI 1640 medium supplemented with 10% Fetal Bovine Serum (FBS), 100 units/ml penicillin and 100 µg/ml streptomycin. We suspended H9 and Sup-T1 cells separately in culture medium at a concentration of 1×10^6 cells/ml, and infected them with HIV for 2 h at a Multiplicity of Infection (MOI) of 0.1. After infection, we then washed the cells twice with PBS and once with the suspension culture medium. The cell suspension (100 µl) was added to each well of a 96-well plate, containing each of the two inhibitors with the concentrations [(0, 1.25, 2.5, 5, 10, 20, 40, and 80 µg/ml)] that equal to (2.3, 4.6, 9.3, 18.6, 37.1, 74.2, 148.4 µM for M522 and 1.7, 3.4, 6.9, 13.9, 27.8, 55.6, 111.2 µM for M32)]. After a four-day incubation, we sub-cultured the cells with fresh culture medium and appropriate concentrations of M522 and M32. We incubated the cells for an additional four days as previously described [15]. Eight days after infection, the antiviral activities of both inhibitors were determined using HIV-1 p24 antigen ELISA [21].

We collected uninfected human blood from a healthy donor and isolated Peripheral Blood Mononuclear Cells (PBMC) by Ficoll-Hypaque density gradient centrifugation. The cells were stimulated using 2 µg/ml of Phytohemagglutinin (PHA) and 32 units/ml of human interleukin 2 (IL-2). We maintained cultures in RPMI 1640 medium supplemented with 15% fetal bovine serum (FBS). We infected cells (5×10^5 cells/ml) with HIV-1RTM or with clinical isolate from HIV-infected patients at 37°C for 2 h. After infection, we washed the cells twice with culture medium, suspended in fresh medium, plated in 24-well plates, and then we added increasing concentrations of M522 inhibitor. After a four-day incubation at 37°C, PBMC cells were sub-cultured in fresh culture medium containing appropriate concentrations of M522 inhibitor and further incubated until day eight after infection. We used the HIV-1 p24 antigen ELISA assay to determine antiviral activities against drug-resistant virus and clinical isolate in PBMC cells. The IC50 ± SD indicated the antiviral activity, where we calculated standard deviation (SD) using KaleidaGraph version 4.0 (Synergy software).

**Disintegration assay**

We obtained a purified recombinant form of the full-length HIV-1 IN (amino acids1–288, wild-type) from the AIDS Research and Reference Reagent Program, Division of AIDS, National Institute of Allergy and Infectious Diseases. Dr. Robert Craigie (Laboratory of Molecular Biology, NIDDK, NIH, Bethesda, MD) generously provided the IN catalytic core domain (amino acids20–212, soluble IN Mutant-type, F185K). We performed the disintegration assay as previously described [20], with the following modifications. A Y-shaped oligonucleotide (i.e., a branched substrate in which the 5′-end was integrated into the target DNA) mimicking the strand transfer (3′-joining) product was used as a substrate. The HIV-1 disintegration substrate was composed of four oligonucleotides (MWG-Biotech, Inc.):

V1/T2 (5′-GACCCCTTTTAGTCAGTGTTGAAAAATCTCTAGCGGTCGACGGCGTCAGTCGAC-3′); T1 (5′-CAGCAACGGCAAGCTTG-3′); T3 (5′TGAGACCTGACGGCCAAAGTCGTGTCGTCG-3′); and USV2 (5′-ACTGCTAGAGATTTCCTCAGACTAAAGGCGGTC-3′). The T1 strand was labelled at the 5′-end with [γ-32P] ATP using T4 polynucleotide kinase as previously described [22]. The disintegration substrate (Y-oligomer) was prepared by annealing the labelled T1 strand with a 3-fold excess of the unlabelled T3, V1/T2, and USV2 strands, incubated at 80°C for 3 min, and then cooled slowly to room temperature. The IN reaction mixture, formulated as previously described [15], was mixed with increasing concentrations of M522 and M32 inhibitors (0, 0.1, 1, 10, 20, 30 and 40 µg/ml). We incubated the HIV-1 IN (300 nM) and the labelled substrate (65 nM) with the reaction mixture for 60 min at 37°C. An equal volume of stop solution
(95% formamide, 30 mM EDTA, 0.1% xylene cyanol, 0.1% bromophenol blue) was added to each reaction and the samples were heated to 95°C for five minutes to denature the DNA. We fractionated the samples by electrophoresis on a 15% denaturing polyacrylamide gel. The substrate was end labelled on the target strand. Disintegration produced a longer DNA product, which generated the conversion of the labelled 16-mer substrate to a 30-nucleotide product. We visualized the product by autoradiography and performed quantitation by phosphorimaging. We then calculated the percent inhibition using the equation: 100 X [(D-C)/(C-N)], where C, N, and D were the fractions of DNA substrate converted to product for DNA alone (C), DNA plus IN (N), and IN plus drug (D). The IC_{50} values of both inhibitors were determined by plotting the log of drug concentrations versus percent inhibition and by determining the concentration that produced 50% inhibition.

**Time-of-drug-addition experiment**

We infected H9 cells with HIV-1\textsubscript{RTMF} at a MOI of 0.5. After 1-h incubation at 37°C, then removed unadsorbed virus by washing three times with RPMI 1640 culture medium. DS, AZT, Ral, M\textsubscript{22} and M\textsubscript{32} were added to the cells at different time points after infection (0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 24, and 25 h) to test their effects on HIV replication, with the exception of DS, which was added before infection (before zero time). The presence of HIV-1 p24 antigen was determined for each inhibitor at 72 h after infection using HIV-1 p24 antigen ELISA assay. We added the reference inhibitors (DS, AZT and Raltegravir) at a standardized concentration corresponding to 100 times their IC\textsubscript{50} values and determined at an MOI of 0.01 as previously described [23].

**Search for emerging of HIV-1 mutants by the selection experiment in H9 cells**

We performed a selection experiment according to a previously described method [24]. We propagated HIV-1\textsubscript{RTMF} in H9 cells in the presence and absence of M\textsubscript{22} and M\textsubscript{32} inhibitors. The cells were infected at a MOI of 0.1 and treated with drug concentrations equal to the 50% inhibitory concentration (IC\textsubscript{50}) of each inhibitor, as determined from the results displayed in Figures 2A and B (4.6 µM for M\textsubscript{22} and 8.3 µM for M\textsubscript{32}). Every three to four days, we sub-cultured the cells with appropriate inhibitor concentrations and then further incubated until day eight after infection. We performed passages by adding the culture supernatant to new H9 cells in the presence of an equal or higher concentration of the inhibitors. Since we observed no cytopathic effects, we sub-cultured the infected cells in the presence of the same concentrations of the inhibitors. The concentrations of inhibitors were gradually increased to 2-, 3-, and 4-times the IC\textsubscript{50} values through the 20 passages of selection (160 days). The antiviral activities were determined in the presence and absence of both inhibitors by using HIV-1 p24 antigen ELISA [21].

**Cytosphere Assay**

We used the assay described by Coulter manual CD4 count kit that contains Coulter CD4 cyto-spheres reagent, inert latex spheres coated with murine monoclonal antibody to identify and manually enumerate by visible light microscopy the absolute count of CD4\textsuperscript{+} cells. We infected H9 cells with HIV-1 in the presence and absence of IN inhibitors (M\textsubscript{22} and M\textsubscript{32}). A comparison of CD4\textsuperscript{+} T-cell counts were determined in three different groups, the first group of cells were infected with HIV-1, the second group of cells were infected and treated with IN inhibitors and the third group were uninfected cells.

**Results**

**Inhibition of acute HIV-1 infection by integrase inhibitors**

To determine the potency of the IN inhibitors M\textsubscript{22} and M\textsubscript{32} (Figure 1), we used different strains of HIV and assayed their replication in H9, Sup-T1 cell lines, and PBMC in the presence and absence of these inhibitors. The antiviral activities of M\textsubscript{22} and M\textsubscript{32} against drug-resistant strains (HIV-1\textsubscript{RTMF}, HIV-1\textsubscript{A17} variant, HIV-1\textsubscript{IIIB/M461I/L69P/V82T/H47Y}, laboratory strains (HIV-1\textsubscript{IIIB} and HIV-1\textsubscript{A17}), and clinical isolate from HIV-infected patients are shown in Figure 2. IC\textsubscript{50} values calculated from the dose response curves are shown in Figures 2A-E. The antiviral activity of M\textsubscript{22} in infected H9 cells against drug-resistant strains was 2.0-11.1 µM while those infected with laboratory strains, was 16.7-20.4 µM (Figure 2A and Table 1). We obtained similar results for M\textsubscript{32}, where the IC\textsubscript{50} values were 1.3-8.3 µM for the drug-resistant strains, and 8.3-9.7 µM for laboratory strains (Figure 2B, Table 1). Experiments on Sup-T1 cells showed similar inhibition of replication for three different strains of HIV-1. The IC\textsubscript{50} value of M\textsubscript{22} against HIV-1\textsubscript{RTMF} was 1.8 ± 1.4 µM, whereas IC\textsubscript{50} for HIV-1\textsubscript{IIIB} was 2.7 ± 1.4 µM and HIV-1\textsubscript{A17} was 2.7 ± 1.1 µM (Figure 2C). M\textsubscript{32} inhibited the HIV-1\textsubscript{RTMF} strain with an IC\textsubscript{50} value of 4.8 ± 0.9 µM, HIV-1\textsubscript{IIIB} with a value of 3.4 µM ± 0.3, and HIV-1\textsubscript{A17} with a value of 2.5 µM ± 0.3 (Figure 2D). In addition, the IC\textsubscript{50} values of M\textsubscript{22} against the replication of HIV-1\textsubscript{RTMF} and clinical isolate in PBMC were 3.5 ± 0.9 µM and 18.0 ± 0.6 µM, respectively (Figure 2E). Both M\textsubscript{22} and M\textsubscript{32} inhibitors exhibited high efficacy against different HIV strains, including clinical isolate from HIV-infected patients and drug-resistant strains, in various cell types (H9, Sup-T1, and PBMC).

**Antiviral activity of IN inhibitors against raltegravir-resistant and wild-type strains**

H9 cells infected with raltegravir-resistant virus harboring double mutations of HIV-1 IN (G140S and Q148H) were treated with different concentrations of the IN inhibitors M\textsubscript{22} and M\textsubscript{32}. We observed sensitivity of the raltegravir-resistant virus to these IN inhibitors. M\textsubscript{22} and M\textsubscript{32} showed high efficacy for inhibiting the replication of this mutant virus with IC\textsubscript{50} values of 1.2 µM ± 0.4 and 2.0 ± 0.7 µM respectively, whereas, the other two control integrase inhibitors, raltegravir (Ral) and IN inhibitor 118-D-24 showed only weak inhibition of the raltegravir-resistant virus (Figure 3A).

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Search for emerging of HIV-1 mutants by the selection experiment in H9 cells

We performed a selection experiment according to a previously described method [24]. We propagated HIV-1,\textsubscript{RTMF} in H9 cells in the presence and absence of M\textsubscript{22} and M\textsubscript{32} inhibitors. The cells were infected at a MOI of 0.1 and treated with drug concentrations equal to the 50% inhibitory concentration (IC\textsubscript{50}) of each inhibitor, as determined from the results displayed in Figures 2A and B (4.6 µM for M\textsubscript{22} and 8.3 µM for M\textsubscript{32}). Every three to four days, we sub-cultured the cells with appropriate inhibitor concentrations and then further incubated until day eight after infection. We performed passages by adding the culture supernatant to new H9 cells in the presence of an equal or higher concentration of the inhibitors. Since we observed no cytopathic effects, we sub-cultured the infected cells in the presence of the same concentrations of the inhibitors. The concentrations of inhibitors were gradually increased to 2-, 3-, and 4-times the IC\textsubscript{50} values through the 20 passages of selection (160 days). The antiviral activities were determined in the presence and absence of both inhibitors by using HIV-1 p24 antigen ELISA [21].
replication of wild-type strain (HIV-1NL4.3) was also tested by both inhibitors and we found that this virus was strongly inhibited with IC50 values of 1.1 µM ± 0.4 and 1.7 ± 0.7 µM, respectively (Figure 3B). Results of replication inhibition for both raltegravir-resistant and wild-type viruses by M22 and M32 are similar to each other.

Figure 2: Effect of M22 and M32 inhibitors on the replication of drug-resistant strains, laboratory strains, and clinical isolate of HIV-1 in H9, Sup-T1, and PBMC cells. Three drug-resistant and two laboratory strains of HIV-1 were treated with increasing concentrations of M22 (Panel A) and M32 (Panel B) in H9 cells. Sup-T1 cells were infected with three strains of HIV-1 (drug-resistant and laboratory strains), and these cells were treated with M22 (Panel C) and M32 (Panel D). PBMC were infected with drug-resistant and clinical isolate from HIV-infected patients and tested with M22 inhibitor (Panel E). The drug-resistant strains are HIV-1RTMF (AZT-resistant), HIV-1L10I/M461/L63P/V82T/184V (Resistant to structurally diverse PR inhibitors), and the laboratory strains are HIV-1L21V and HIV-1MN. Viral replication in the absence and presence of different drug concentrations and the IC50 values were determined from the dose response curve. The results obtained from the means and standard deviations from three independent experiments performed in duplicate.

Interaction of M22 and M32 with HIV-1 Integrase

The effects of both inhibitors were tested against the catalytic core domain (IN50-212, mutant-type) as well as the full-length enzyme (IN1-288, wild-type) using the disintegration assay to determine whether M22 and M32 are able to block HIV-1 IN activity. M22 inhibited the disintegration activity catalyzed by the catalytic core (IN50-212) with an IC50 value of 4.0 µM as illustrated in Figures 4A and 4C, and inhibited the full-length enzyme (IN1-288) with an IC50 value of 3.3 µM (Figures 4B and 4C). Similarly, M32 inhibited the activity of the catalytic core domain (IN50-212) with an IC50 value of 6.2 µM (Figures 4D and 4F), and the full-length IN (IN1-288) with an IC50 value of 5.8 µM (Figures 4E and 4F). Overall, these results indicated that both M22 and M32 inhibited the activity of IN50-212, demonstrating that the integrase catalytic core region is the binding site for both inhibitors. The interaction between the inhibitors and the binding site is responsible for prevention of viral integration and inhibition of HIV-1 IN.

Table 1: Antiviral activities of M22 and M32 against drug-resistant, laboratory strains of HIV-1 in H9 cells. Anti-HIV activity (IC50), 50% inhibitory concentration or concentration of each inhibitor required to inhibit viral replication by 50 % and calculated as (µM) ± SD. Results were obtained from three independent experiments performed in duplicate.

Figure 3: Inhibition of the replication of raltegravir-resistant and wild-type strains in H9 cells by M22 and M32 inhibitors. Cells were infected with raltegravir-resistant virus that contains the integrase mutations G140S and Q148H. The infected cells were treated with increasing concentrations of M22 (Panel A) and M32 (Panel B) in H9 cells.
Figure 4: Inhibition of disintegration activity of catalytic core domain and full-length HIV-1 integrase by M522 and M532. The disintegration assay measured the conversion by HIV-1 IN of the radiolabeled 16-base oligonucleotide of the Y-shaped substrate (see Methods) to a 30-base disintegration product. Increasing concentrations (0, 0.1, 1, 10, 20, 30, and 40 µg/ml) of M522 (Panels A and B) or M532 (Panels D and E) were added to disintegration assay reactions catalyzed by either the catalytic core domain (IN50-212) or full-length HIV-1 integrase (IN1-288). A graphic representation of the data for the disintegration assays (Panels A, B, D, and E) as quantitated by phosphorimaging was presented, showing a dose-response inhibition of disintegration activity by M522 (Panel C) and M532 (Panel F) for the full-length HIV-1 integrase (Wild-type) and its catalytic core domain (Mutant-type).

Time of intervention on HIV replication cycle

To investigate which step of the replication cycle was inhibited by M522 and M532, we performed a time-of-drug-addition experiment. In addition, we also evaluated when these IN inhibitors were needed in the viral replication cycle and for examining the maximum time that is permissive before these inhibitors lose their antiviral efficacy. We infected H9 cells at a high MOI (0.5) to synchronize all steps of viral replication. Three reference inhibitors, dextran sulfate (DS), AZT, and raltegravir (Ral), with known modes of action were used as controls for comparison with M522 and M532. Dextran sulfate (DS) is known to interfere with binding of the virus to the cell, the nucleoside analogue AZT inhibits the reverse transcription process, and raltegravir (Ral) is known to interfere with viral integration. Our data clearly indicated that addition of DS to the H9 cells before the viral infection was necessary, because its antiviral activity was lost when it was added after viral infection (Figure 5A). Conversely, the addition of AZT can be delayed up to 4 h after infection (Figure 5B), while raltegravir (Ral) only lost its activity when added more than 8 h after infection. IN inhibitors M522 and M532 produced a clearly different profile than the DS and AZT inhibitors (Figures 5A and 5B), while sharing an identical profile with that of the Ral inhibitor (Figure 5C). These results strongly suggested that M522 and M532 did not inhibit viral entry and reverse transcription, but they interfered with the viral replication cycle at a point coinciding with the integration step.

Figure 5: Effect of time of drug intervention on HIV replication cycle. H9 cells were infected with HIV-1 at a MOI of 0.5 and the test inhibitors were added at different times after infection (Panels A, B, and C). Viral p24 antigen values were determined at 72 h after infection for the means of three independent experiments and error bars represent standard deviations. The reference inhibitors DS (a viral binding inhibitor, Panel A); AZT (RT inhibitor, Panel B); and Ral (an IN inhibitor, Panel C) were used in parallel and compared with the M522 and M532 inhibitors versus the control. Arrows denote time points that the inhibitors lost their antiviral activity after this time.

Search for emerging of HIV-1 mutants by the selection experiment in H9 cells

To evaluate the propensity of these inhibitors to induce new resistance, we propagated HIV-1RTMF in H9 cells with the constant presence and absence of M522 and M532 inhibitors. These IN inhibitors were added at their IC50 values in the first passage of virus and the culture supernatant from each infection was used to infect new cells. We serially passed the virus with increasing concentrations of both inhibitors to 2-, 3-, and 4-times the IC50 values during the selection experiment. The antiviral activities of both inhibitors were determined against the selected strain in H9 cells for each passage. The results demonstrated that the selected virus was unable to replicate and no drug-resistant mutants developed during the passage of virus in the presence of M522 and M532 over 20 passages (160 days) versus the control virus without inhibitors, indicating that the HIV-1RTMF remained sensitive to both inhibitors even after 20 viral passages (Figure 6A).
Syncytia are giant multinucleated structures, clusters of many nuclei within a single cell membrane, and are a sign of HIV infection in cell cultures. They formed when HIV-infected cells make gp120 and carry it on their surface, fused with healthy cells bearing CD4+ molecules (Figure 6B). The second group of infected cells was treated with the IN inhibitors, where they inhibited HIV-1 replication, prevented syncytia formation and protected the cells from cytopathic effects (Figure 6C). The third group was uninfected H9 cells as a negative control to be compared with cells infected with HIV-1, treated and non-treated cells with the IN inhibitors. The treated cells with M22 and M32 were found to be morphologically identical to the uninfected cells (Figure 6D).

In addition, we performed more studies to test the protection effect of these IN inhibitors. A comparison of CD4+ T-cell counts was done in four different groups using cytosphere assay. We infected two groups of H9 cells with HIV-1 (AZT-resistant) and treated with the inhibitors (M22 and M32), the third group was infected but not treated as a positive control, and the fourth group was uninfected cells as a negative control. The absolute CD4+ T-cell counts were calculated for all four groups, where the number of CD4+ T-count for the infected cells with HIV and treated with both inhibitors was very low at day zero then increased and back to the normal level as found for the uninfected cells after 8 days. Therefore, the groups treated with both IN inhibitors and the uninfected group showed an identical pattern and had higher CD4+ counts per µl than the group of cells that was infected but not treated after 8 days (Figure 6E).

Furthermore, a comparison for quantitation of p24 antigen between the treated and non-treated groups of the infected cells with both inhibitors versus the uninfected cells was also calculated. The cells infected with HIV-1 and treated with M22 and M32 did not show any p24 antigen and are identical to the uninfected cells as if they were not infected, whereas the cells infected but non-treated had very high p24 antigen after 8 days (Figure 6F).

## Discussion

In the current study, we evaluated the antiviral efficacies of the IN inhibitors (M22 and M32) against wild-type virus, different drug-resistant HIV-1 strains and clinical isolate from infected patients. We found M22 and M32 are effective in stopping viral replication of many mutant strains that are resistant to RT, IN and PR drugs. By stopping the viral infection, M22 and M32 protected the infected cells from cytopathic effects caused by the virus yielding healthy T cells with high CD4+ cell counts identical to those of uninfected cells.

FDA approved raltegravir as an HIV-1 IN inhibitor in treatment-experienced patients [26] and resistance to raltegravir in vivo was previously described [4]. This resistance due to the mutation combination of G140S/Q148H which is most relevant in vivo with the highest increase in resistance factor [27-29], and exhibits the greatest loss of drug susceptibility [30]. Therefore, the emergence of drug-resistant HIV variants can attenuate the efficacy of antiretroviral treatment [31]. Our IN inhibitors M22 and M32 effectively suppressed the replication of resistant viruses containing G140S/Q148H mutation that was isolated from patients in whom raltegravir treatment had been ineffective. Moreover, M22 and M32 have higher efficacies in inhibiting different drug-resistant HIV-1 strains and clinical isolate when compared to the scutellarin inhibitors [32], as well as having higher efficacies than the IN inhibitors of L-chicoric acid analogues for the drug-resistant clone [22]. Furthermore, our IN inhibitors M22 and M32 have higher efficacies in inhibiting different drug-resistant HIV-1 strains and clinical isolate when compared to the scutellarin inhibitors [32], as well as having higher efficacies than the IN inhibitors of L-chicoric acid analogues for the drug-resistant clone [22]. Furthermore, our IN inhibitory efficacies on infected cells with HIV-1.

### Protective effect of M22 and M32 on infected cells with HIV-1

To examine the protective effect of the IN inhibitors (M22 and M32) on CD4+ T-cells, two groups of H9 cells were infected with HIV-1RTMF (AZT-resistant). The first group of the infected cells was in the absence of the IN inhibitors, and these cells formed syncytia. The absolute CD4+ T-cell counts were calculated for all four groups (Panel F) where the values obtained from the means and standard deviations of three independent experiments.
inhibitors showed higher activities against HIV-1 strains resistant to non-nucleoside RT (NNRTI), than the IN compounds L-708,906 and V-165 [33]. In addition, M22 and M32 have high viral suppression against raltegravir-resistant and wild-type viruses showing a comparable pattern as displayed in their IC_{50} values. Overall, M22 and M32 inhibitors possess high potency against wild-type virus and different drug-resistant HIV-1 strains, particularly the clinically relevant integrase-resistant one of raltegravir.

The disintegration assay provided an effective tool for measurement of drug effects on IN deletion mutants [19]. We examined the interaction of M22 and M32 inhibitors by testing them with the catalytic core domain of HIV-1 IN and full-length IN. Both IN inhibitors showed higher efficiencies against disintegration activity than those of other IN inhibitors; integric acid, equisetin [34], and α-hydroxytropolone [35]. On comparison to the IN inhibitors, raltegravir and elvitegravir, M22 and M32 also showed high inhibition of disintegration activity, while both raltegravir and elvitegravir showed weak or no inhibition of this reaction [36]. Overall, we found that M22 and M32 inhibited disintegration activity by interacting with the catalytic core region and inhibit HIV-1 IN in a manner consistent with that of other IN inhibitors [37].

We have previously shown that M22 and M32 inhibited the activities of HIV-1 IN, 3’-processing and 3’-joining to the target DNA (strand transfer) [15]. Our current data also demonstrated that both M22 and M32 inhibited the disintegration activity with equivalent potency in the full-length IN and in its catalytic core domain. This is indicating that the catalytic core region is the target of interaction for both inhibitors and is responsible for the inhibition of HIV-1 IN and prevention of viral integration. Although M22 and M32 inhibitors bind to the catalytic core domain like some other FDA approved drugs of INSTIs, the potency of these inhibitors must come from their chemical structures; which play a potential role for binding interactions with HIV IN. Structural features of these two inhibitors that are crucial for the enzymatic activity are hydroxy, aromatic ring and carboxylic acid moieties. To clarify potency of these two IN inhibitors, a recent reported study of molecular docking was conducted to determine the binding modes of M22 and M32 inhibitors within the catalytic core site of HIV-1 IN by using PFV IN model for WT (wild-type) and mutant variant, S217H mutant, S217H mutant (equivalent to G140S/Q148H HIV-1 IN). The docking calculations showed that the major key interactions for stabilization of M22 and M32 binding are hydrogen bonding, metal-ligand, and π-π stacking interactions with adenosine base A17 of the viral DNA, while these key interactions were not observed with raltegravir [38]. Another molecular docking study was conducted for M22 and M32 inhibitors using of HIV-1 IN [39] instead of PFV IN [38]. The docking results with HIV-1 IN indicated that the binding modes of M22 and M32 were similar to those of 5-CITEP inhibitor [39]. Moreover, the docking calculations showed that the carboxyl and hydroxyl groups on the side-chain of M32 are important chemical groups, which make M32 a tighter binder than M22 [39]. Overall, these two docking studies are supportive to our findings on the uniqueness of M22 and M32 as strong HIV-1 IN inhibitors.

The CD4+ cell count at initiation is the prognosis factor of HIV-1 in infected patients for a highly active antiretroviral therapy (HAART) [40]. In this case, viral load and CD4 cell counts serve as the two surrogate markers for monitoring of antiretroviral treatment (ART) responses and HIV disease progression where measurement of CD4 count is particularly useful before initiation of ART [41]. For these reasons, we have used the effect of M22 and M32 treatment on the AZT-resistant virus (HIV-1\textsuperscript{R}\textsubscript{ TAM}) in culture as a mimic to HIV-infected patients. Similar to AIDS patients under an effective ART, although we found the CD4+ counts to be very low in non-treated and in infected cells during the early phase of the treatment, continuing M22 and M32 treatments not only blocked HIV-1\textsuperscript{R}\textsubscript{ TAM} from further infection, they also brought numbers of CD4+ cells back to normal and identical to those of the uninfected cells after 8 days. This is an indication that these IN inhibitors are not toxic to the cells and protected the infected cells with drug-resistant virus from depletion of CD4+ T-cells. All the presented data of the current study show the high inhibition activity of these IN inhibitors against drug-resistant strains and they did not build up any further drug resistance, at least in cell culture.

Conclusion

Our findings showed that the IN inhibitors M22 and M32 strongly suppressed the replication of different of HIV-1 strains (wild-type; different drug-resistant strains; clinical isolate from HIV-infected patients; and laboratory strains) grown in different cell types (H9, Sup-T1, and PBMC). In addition, they critically inhibited the viral replication of the clinically relevant raltegravir-resistant virus and did not induce drug resistance during the selection study. Moreover, disintegration studies support a model in which both inhibitors interacted with the catalytic core domain of HIV-1 IN, blocked its activity and inhibited viral replication. Time of intervention on HIV replication study also indicated that M22 and M32 interfered with the viral replication cycle at a time point coinciding with the integration step. Furthermore, these IN inhibitors protected the infected cells with HIV-1 from cytopathic effects and CD4+ T-cells depletion. This study has highlighted the application of M22 and M32 as candidate IN inhibitors for drug development against diverse HIV-1 strains, particularly for those drug-resistant HIV strains. These two selective IN inhibitors M22 and M32 hold the promise as a novel class of therapeutic drugs for AIDS patients based on their high potentials and in the absence of cytotoxicity.

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