Preclinical and Clinical Profile of HIV-1 Integrase Strand-transfer Inhibitor GS-9224 Compared to its Parent Compound GS-9160

Gregg S Jones, Rebecca Hluhanich, Lani M Wieman, Jim Zheng, Wayne Huang, April Kinkade, Eugene J Eisenberg, Chris Yang, Damian McColl, Anita Mathias, Haolun Jin, Tomas Cihlar, Matthew Wright, Romas Geleziunas and Manuel Tsiang*

Gilead Sciences, Foster City, CA 94404, USA

Abstract

In an effort to optimize the pharmacokinetic profile of GS-9160, a potent (EC50=1.2 – 4 nM) antiviral tri-cyclic HIV-1 integrase (IN) strand transfer inhibitor (INSTI), various substitutions on the p-fluorobenzyl moiety of GS-9160 were explored. This effort led to the discovery of GS-9224, an analog containing a 5-chloro-2,4-di-fluorobenzyl moiety. GS-9224, like its predecessor GS-9160, has potent (EC50=1.3 – 8 nM) and selective antiviral activity against HIV-1 and acts as a bona fide integration inhibitor through elevation of 2-long terminal repeat (2-LTR) circles and decrease of integration junctions in HIV-1 infected cells, markers of failed viral integration. Viral resistance selections with GS-9224 yielded three mutations within the catalytic core domain of HIV-1 IN: G140S, L74M and Q148K. When tested against a panel of HIV-1 clones engineered to express INSTI resistance mutations, the profile of GS-9224 was comparable to GS-9160. GS-9224 exhibits improved hepatic microsome stability, better absorption potential and a lower in vitro intestinal efflux ratio compared to GS-9160. In addition, GS-9224 displayed higher hepatic microsome stability, better absorption potential and serum protein binding of GS-9224 compared to GS-9160 did not translate into an improved clinical pharmacokinetic profile. Results from a single-dose study in human healthy subjects revealed a fast systemic clearance of GS-9224 with a terminal half-life of ~ 1 h, resulting in a pharmacokinetic profile that would not support once-daily dosing in HIV-infected patients.

Keywords: HIV-1; GS-9224; GS-9160; Strand-transfer

Introduction

As an essential enzyme for HIV-1 replication, integrase (IN) is an attractive target for antiviral drug discovery. During the viral life cycle, the primary function of IN is to catalyze the insertion of reverse-transcribed double stranded HIV-1 DNA into the host cell genome to establish the provirus. Inhibition of the strand transfer activity of IN as a target for therapeutic intervention is now well validated clinically [1,2]. Raltegravir (RAL) was the first IN strand-transfer inhibitor (INSTI) to be approved for clinical use and is dosed twice daily [3-5]. Elvitegravir (EVG) is co-administered with cobicistat, a new pharmacokinetic enhancer that enables once-daily dosing of EVG [6-8]. A once-daily single tablet regimen (STRIBRILD™) containing EVG, cobicistat, deoxyfluorothiacytidine (FTC) and tenofovir disoproxil fumarate (TDF) has been recently approved by the FDA for the treatment of HIV infection. In two Phase III studies, the STRIBILD regimen showed non-inferior efficacy and safety in treatment-naïve HIV-infected patients when compared head-to-head with either efavirenz [9] or atazanavir [10] containing once-daily treatment regimens. Recent data from the 96-week extension analysis of both studies established the longer term durability and efficacy of the STRIBILD regimen [11,12]. Specific mutations in IN conferring cross-resistance to RAL and EVG have been identified in clinical settings [13-15]. The INSTI dolutegravir (S/GSK1349572), which is dosed once-daily without boosting, displayed a resistance profile that is distinct from those of RAL and EVG [16,17]. In phase III clinical studies, none of the subjects treated with the combination of dolutegravir and TDF/FTC for up to 48 weeks developed any treatment-associated IN mutations [18].

In an effort to identify additional novel INSTIs that could be dosed once-daily without pharmacokinetic enhancement we explored modifications of a tricyclic analog GS-9160, a potent and orally bioavailable compound [19-21]. Substitutions on the p-fluorobenzyl moiety of GS-9160 led to the discovery of GS-9224, which contains a 5-chloro-2,4-di-fluorobenzyl moiety [22]. In this report, we describe detailed biological and metabolic characterization of GS-9224 including a clinical pharmacokinetic profile in healthy human volunteers.

Materials and Methods

Compounds

GS-9224 and GS-9160 were synthesized at Gilead Sciences, Inc. L-870,810 was synthesized by Combi Blocks, Inc. (San Diego, CA). RAL was synthesized by Naeja Pharmaceuticals, Inc. (Edmonton, Alberta Canada). All three compounds have a purity of >95% as determined by analytical reversed phase HPLC using a Phenomenex Luna C18 column and UV detection at 214 nm and 254 nm. The structures of these three INSTIs are shown in (Figure 1).

Cells

MT-2 cells were obtained from Stanford University and MT-4 cells were obtained from the NIH AIDS Research and Reference Reagent Program (Germantown, MD). SupT1 cells were obtained from the American Type Culture Collection (Rockville, MD). MT-2, MT-4 and SupT1 cells were maintained in RPMI-1640 medium (Cat # 61870-

*Corresponding author: Manuel Tsiang, Gilead Sciences, 333 Lakeside Drive, Foster City, CA 94404, USA, Tel: (650) 522-5860; Fax: (650) 522-5143; E-mail: Mtsiang@gilead.com

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To remove un-joined target DNA, and the chemiluminescence signal was detected following addition of horseradish peroxidase-conjugated anti-digoxigenin antibody. Test compounds were added immediately before the target DNA.

**Antiviral and cytotoxicity assays**

Antiviral and cytotoxicity assays in HIV-1 IIib infected MT-2, MT-4 and SupT1 cells and PHA plus IL-2 stimulated human PBMCs have been described in the materials and methods section of an earlier study by our investigators at Gilead Sciences [19]. Briefly, for the antiviral assay utilizing MT-2 and MT-4 cells, 50 μL of 2X test concentration (total of 9 concentrations) of 3-fold serially diluted compound in culture medium with 10% FBS was added to each well of a 96-well plate in triplicate. MT-2 cells were infected with HIV-1 IIb virus at a multiplicity of infection (MOI) of ~0.01 for 3 hours at 37°C. Fifty microliters of infected cell suspension in culture medium with 10% FBS (~1.5 x 10^6 cells) was then added to each well containing 50 μL of diluted compound. The plates were then incubated at 37°C for 5 days. At the end of the incubation, 100 μL of CellTiter-Glo™ Reagent (catalog # G7571, Promega Biosciences, Inc., Madison, WI) was added to each well. Cell lysis was carried out by incubating the assay plates at room temperature for 10 min followed by the quantification of chemiluminescence signal using the Envision plate reader (Perkin Elmer, Shelton, CT). For compound cytotoxicity assessment in MT-2 and MT-4 cells, a protocol identical to that of the antiviral assay was followed, except that uninfected cells were used and a higher concentration range of compound was tested.

To study the effect of serum proteins on the antiviral activity of INSTI, compounds were tested in the presence of either 35 mg/ml HSA (catalog # A-1653, Sigma, St. Louis, MO), or 1.5 mg/ml α-Amylase (catalog #G-9885, Sigma, St. Louis, MO). To test the effect of human serum on compound potency, the assay was performed in the presence of 10%, 20%, 35% and 50% human serum from clotted whole blood (catalog #14-498E, Cambrex Bio Science, Walkersville, MD).

**2-LTR circle accumulation assay**

The assay for 2-LTR circle accumulation was adapted from a previously described method [26]. SupT1 cells were infected in bulk culture with HIV-1 IIIB at a multiplicity of infection (MOI) of 10 and at a cell density of 4 x 10^6 cells/ml for 3 hours at 37°C. We have confirmed in a time course experiment in our own system a previous observation that 2-LTR circles reached a peak production 24 hours post-infection [26]. For this reason, measurement of 2-LTR circles was performed at 24 hours post-infection in our inhibition dose response studies. Both the 2-LTR junction (2-LTR circles) and the host globin gene (used to normalize for cell number) were quantified using Taqman real-time PCR [19]. The copy number of 2-LTR circles was normalized to that of the globin gene and plotted against drug concentration for EC_{50} determination.

036, Life Technologies, Grand Island, NY) supplemented with 10% heat-inactivated fetal bovine serum (FBS) (Cat # SH30071.03, GE Healthcare Sciences, Logan, UT) and antibiotics. Peripheral blood monocytes (PBMCs) were isolated from the buffy coats of healthy human volunteers (Stanford Medical School Blood Center, Stanford, CA).

**Viruses**

HIV-1 strain IIIb was obtained from Advanced Biotechnologies Inc. (Columbia, MD). The NRTI-resistant virus containing mutation K65R in reverse transcriptase (RT) was obtained from Mark Wainberg (McGill AIDS Center, Montreal, Canada). The NRTI-resistant virus RT-M184V and NNRTI-resistant viruses RT-K103N and RT-Y181C were constructed by site-directed mutagenesis [23]. The NRTI-resistant virus, RT-6Tam (M41L, D67N, K70R, L210W, T215Y and K219Q) which confers resistance to thymidine analogs [23] as well as the protease (PR) inhibitor-resistant viruses, PR-I84V/L90M and PR-G48V/V82A/L90M were constructed by homologous recombination in electroporated SupT1 cells [24]. All the above mutant viruses were constructed from an HXB2 clone.

**Construction of infectious HIV-1 DNA clones with IN mutations**

Mutations conferring resistance to INSTI were introduced into the infectious HIV-1 DNA clone HXB2 by site-directed mutagenesis at Gilead Sciences by our investigators [19]. Briefly, the 15.5 Kb HXB2 plasmid DNA was cut with ApaI and SalI to generate a 3.7 Kb ApaI – SalI fragment containing the mutation(s) by sequencing, DNAs were prepared from the cultures, and the ApaI - SalI fragment that was gel-purified. This 3.7 Kb fragment containing the wild-type 3.7 Kb ApaI – SalI fragment was then cloned into pUNI/V5-His B (Invitrogen, Carlsbad, CA) to generate pHXB2 (3.7). After confirmation of the mutation(s) by sequencing, both the RT and the IN coding regions was then subcloned into an HXB2 clone.
**Alu-PCR Assays**

The Alu-PCR assay has been previously described [26]. MT-2 cells were infected as a bulk culture with pseudotyped HIV-1 vector particles produced from the SOD1k 2G cell line [26,27] at an m.o.i. of 10 and a cell density of 4 x 10⁶ cells/ml for 3 hours at 37°C. The plates were incubated at 37°C for 12 hours for Late-RT product quantification and for 48 hours for Alu-PCR product quantification. The levels of Late-RT sequences and integration junctions quantified at 12 and 48 hours post-infection respectively were normalized to the level of globin gene present in the sample at these time points. Quantification by PCR was performed on the ABI Prism® 7900HT Sequence Detection System (Applied Biosystems, Foster City, CA).

**Data analysis**

The entire curve fitting to generate EC₅₀ and CC₅₀ as well as error calculations were performed using Sigma Plot scientific data analysis and graphing software (Systat Software Inc., San Jose, CA).

**Viral resistance selection**

Resistance selections were carried out in 6-well tissue culture plates. MT-2 cells were seeded at a density of 0.5 x 10⁶ cells per well in 5 ml of culture medium. Compounds were added at a final concentration corresponding to their antiviral EC₅₀ or twice the EC₅₀. HIV-1 IIIb was used at an MOI of ~0.01. The cultures were incubated at 37°C and split 1/2-1/3 once or twice a week depending on the growth status of the cells. Cytotoxic effect (CPE) manifested as syncytia formation was used to follow progression of infections. Virus was harvested and transferred to fresh MT-2 cells in the presence of higher concentrations of compounds. Successive viral passages were obtained by repeating this procedure. The duration of each passage ranged from 10-15 days.

**Population and clonal sequencing of HIV-1**

Viral RNA isolated from culture supernatants was used for cDNA synthesis. Five µl of cDNA was used for PCR amplification of a 2765 base pair fragment spanning nucleotide 372 of the RT coding region to nucleotide 127 of the vpr gene and encompassing the entire IN open reading frame. The following primers were used for PCR: pLAIR3004: 5’-CAGGAAGTATACTGCAGTTACC-3’ and pLAIR5768: 5’-CTAAGCGCATGGACGAAATC-3’ (BioSource International, Foster City, CA). The PCR product was purified using the QIAquick PCR purification kit (Catalog #28104, QIAGEN, Valencia, CA). This PCR product was used for population sequencing of drug-resistant viral pools and both DNA strands were sequenced (ELIM Biopharmaceuticals Inc, Hayward, CA). For clonal sequencing, a 1296 base pair PCR product spanning nucleotide 1608 of the RT coding region to nucleotide 372 of the RT coding region was used as template. The PCR product was purified on a 1.0% agarose gel and cloned into the pCR-Blunt II-TOPO plasmid using the Zero Blunt TOPO PCR Cloning Kit (Catalog #K2800-20, Invitrogen, Carlsbad, CA). Thirty transformed bacterial colonies were picked and cultured overnight in L-Broth containing 50 µg/ml kanamycin. Plasmid DNAs were isolated using a Wizard SV 96 Plasmid purification kit (Catalog #A2255, Promega, Madison WI) and sequenced through the IN region on both DNA strands (ELIM Biopharmaceuticals Inc, Hayward, CA) using the following primers: INTseq14157: 5’-ACACACACACAACAAAGGAAATGG-3’ and VIFseq2Rev5453: 5’-CTGGCTTGTATTTCCACCC-3’ (BioSource International, Foster City, CA).

**Hepatic microsomal stability assay**

Microsome (MS) suspensions were prepared by dilution of stock microsomes (In Vitro Technologies, MD) with 0.05 M phosphate buffer, pH 7.4 to obtain protein concentrations of 0.5 mg/mL. Compounds were mixed with MS suspension media (pre-heated for 5 min at 37°C) and cofactors were added to start the reaction. The final reaction mixture contained: 2 µM compound, 0.5 mg/mL microsomal protein, 1.25 mM NADPH, 3.3 mM MgCl₂ and 0.05 M phosphate buffer, pH 7.4. At 0, 5, 15, 30, 45 and 60 min, 25 µL of the reaction mixture were quenched with 250 µL of internal standard (50 nM)/mobile phase B. The analytical internal standard was suspended in 0.2% formic acid/95% acetonitrile/5% water. After quenching, the plates were centrifuged at 3000 × g for 10 min. Five µL samples were analyzed by LC/MS/MS (LOQ = 0.001µM). Verapamil was used as a positive control for microsomal activity.

**Caco-2 permeability**

Caco-2 cells were grown to confluence on 24 well PET (polyethylene-terephthalate) plates (BD Biosciences, San Jose, CA). Donor HBSS buffer from Invitrogen was supplemented with 10 mM HEPES and 15 mM glucose with pH adjusted to pH 6.5. Receiver HBSS buffer was supplemented with 1% BSA with pH adjusted to pH 7.4. After equilibration with transport buffer, trans-epithelial electrical resistance was measured to confirm membrane integrity. The assay was initiated by the addition of compound containing buffer to the donor compartment. Samples of 100 µL were removed from the receiver compartment at 1 and 2 h post compound addition. The removed samples were immediately mixed with 400 µL of 0.1% formic acid in acetonitrile to precipitate protein and stabilize test compounds. Cells were dosed on either the apical or baso-lateral side to determine the forward (A to B) and reverse (B to A) permeability. Permeability through the membrane and non-specific binding was measured through a cell free trans-well. To test for non-specific binding and compound instability, the total amount of compound was quantified at the end of the experiment and compared to the material present in the original dosing solution to determine % recovery. Samples were analyzed by LC/MS/MS.

**Plasma protein binding assay**

Equilibrium dialysis was conducted at 37°C using Beagle dog and human plasma containing either GS-9224 or GS-9160 at the final concentrations of 2 µM. Based on a pilot time-to-equilibrium study and stability data, a 30-hour dialysis time was chosen for plasma protein binding determinations. Prior to the study, dialysis membranes were soaked for approximately one hour in 0.133 M phosphate buffer, pH 7.4. Plasma samples containing compound (1 mL) and compound-free phosphate buffer (1 mL) were placed into opposite sides of the assembled dialysis cells. The dialysis cells were rotated slowly in a 37°C water bath for 30 h. Plasma samples were next drained into preweighed polypropylene tubes containing 1 mL of phosphate buffer. Buffer samples were drained into weighed tubes containing 1.0 mL of blank plasma of the matching species. Post-dialysis plasma and buffer weights were measured and recorded for calculations.

**Pharmacokinetic profiling in healthy human subjects**

Ten healthy subjects were randomized to receive a single dose of GS-9224 (n=8) or matching placebo (n=2) following an overnight fast. Doses of 50 and 100 mg were administered in orange juice on day 1 and day 9 to the same patients (including a rinse of the dispensing container) to a total liquid volume of 240 mL with dosing within 15 minutes of study drug preparation. Following study drug administration, subjects were restricted from water intake until after the 2-hour blood draw and
from food intake until after collection of the 4-hour blood draw. Serial blood samples were collected at times: 0 (pre-dose), 0.25, 0.5, 0.75, 1, 1.5, 2, 2.5, 3, 3.5, 4, 5, 6, 8, 10, 12, 18, 24, 28, 32, 36, 40, 44, and 48 hours postdose. In subjects that received GS-9224 plasma concentrations were measured at all-time points; protein binding of GS-9224 was assessed at the 1-hour time point.

Assessment of adverse events and concomitant medications continued throughout the duration of the study. Laboratory analysis, serial ECGs, and physical examinations were performed at screening, at defined intervals during the confinement periods, and at the follow-up evaluation 7 days after the last dose of study drug administration.

The study protocol and informed consent document(s) were reviewed and approved by the study center’s Institutional Review Board and subjects were provided written informed consent before study participation. The studies were carried out in accordance with the clinical research guidelines established by the Basic Principles defined in the U.S. 21 CFR Part 312.20 and the principles enunciated in the latest version of the Declaration of Helsinki.

### Results

**GS-9224 exhibits potent antiviral activity and is synergistic with HIV antiviral drugs**

In an effort to improve the pharmacokinetic profile of previously published INSTI GS-9160 [19-21], various substitutions on the p-fluorobenzyl moiety of GS-9160 were explored leading to the discovery of the close analog GS-9224 containing a 5-chloro-2,4-fluorobenzyl moiety (Figure 1) [22].

The antiviral activity of GS-9224 was evaluated in HIV-1 infected MT-2 and MT-4 cell lines as well as in primary human PBMCs (Table 1). The anti-HIV activity of GS-9224 was similar in all three cell types with an EC_{50} range of 1.3 to 9 nM. This potency is within a range similar to that observed for RAL and slightly lower than the parent analog GS-9160. To assess the impact of human serum protein binding on the antiviral potency of GS-9224 and to provide a target trough concentration in humans to achieve efficacy, the antiviral activity of GS-9224 was determined in the presence of human serum and the human serum components including human serum albumin (HSA) and α1-acid glycoprotein (α1-AGP) (Table 2). The EC_{50} of GS-9224 in 100% human serum was extrapolated from the EC_{50} values determined in the presence of 10, 20, 35 and 50% human serum. The combined effect of both serum proteins, HSA and α1-AGP increased the EC_{50} of GS-9224 by ~14.6-fold in MT-2 cells, while the EC_{50} values of GS-9160 and RAL were increased by 10.8- and 3-fold, respectively. Whole human serum had an effect similar to the combined serum proteins in MT-2 cells and increased the EC_{50} of GS-9224 by 12.3-fold. In human PBMCs, the combined effect of HSA and α1-AGP increased the EC_{50} of GS-9224 by 8.6-fold, a similar shift in the EC_{50} values as that observed for GS-9160 and RAL (10.8- and 7.3-fold, respectively).

Similar to GS-9160, GS-9224 showed synergistic antiviral effect in HIV-1-infected MT-2 cells when combined with approved drugs from different antiretroviral classes including PIs (lopinavir, atazanavir, and nelfinavir), NNRTIs (efavirenz) and N(t)RTIs (tenofovir, zidovudine, emtricitabine, and lamivudine) (Table S1).

**GS-9224 inhibits HIV-1 integrase function and integration in infected cells**

The inhibitory effect of GS-9224 on IN enzymatic activity was evaluated in a biochemical assay that measures the activity of IN in an *in vitro* integration reaction mediating the strand transfer of viral donor DNA unto host or target DNA. GS-9224 produced inhibitory activity (IC_{50} = 55 nM) that was comparable to that of GS-9160 (IC_{50} = 29 nM) and RAL (IC_{50} = 34 nM) (Table 3). An elevation in the levels of circular HIV DNA species containing 1- or 2-long terminal repeats (1-LTR and 2-LTR circles) following treatment of infected cells with a small molecule can provide evidence that HIV integration has been inhibited [28]. Treatment of HIV-1 infected SupT1 cells with GS-9224 caused a ~3.9-fold increase in 2-LTR circles. The effect was dose-dependent with an EC_{50} of 9.5 nM (Table 3). Similar results were obtained with the parental analog of GS-9224, GS-9160 and the clinically validated integrase strand-transfer inhibitor, RAL. This result provided initial evidence that GS-9160 can block HIV-1 integration in a cell.

Another method to assess whether HIV-1 integration is inhibited in infected cells is by direct measurement of integration junctions between HIV proviral DNA and the host chromosomal DNA [26]. Detection by PCR of nucleic acid products containing Alu-repeat sequences

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### Table 1: Antiviral Activity of GS-9224 in Various HIV-1 Infected Human Cell Types.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Anti-HIV in MT2</th>
<th>Cytotoxicity in MT2</th>
<th>Anti-HIV in MT4</th>
<th>Cytotoxicity in MT4</th>
<th>Anti-HIV in PBMC</th>
<th>Cytotoxicity in PBMC</th>
</tr>
</thead>
<tbody>
<tr>
<td>GS-9224</td>
<td>1.3 ± 1.1</td>
<td>1600 ± 600</td>
<td>9.4 ± 0.5</td>
<td>700 ± 300</td>
<td>5 ± 6</td>
<td>5850 ± 900</td>
</tr>
<tr>
<td>GS-9160</td>
<td>1.2 ± 0.5</td>
<td>4200 ± 1000</td>
<td>4 ± 3</td>
<td>800 ± 400</td>
<td>1.2 ± 0.9</td>
<td>3880 ± 1700</td>
</tr>
<tr>
<td>RAL</td>
<td>7.5 ± 1</td>
<td>&gt;100,000</td>
<td>8 ± 2</td>
<td>58200 ± 7300</td>
<td>2.6 ± 1.2</td>
<td>210.500 ± 90,000</td>
</tr>
</tbody>
</table>

a Mean ± SD of at least 4 independent experiments.
b Mean ± SD of at least 3 different donors.

### Table 2: Effect of Serum Proteins on the Antiviral Activity of GS-9224.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Anti-HIV in MT2 Cells</th>
<th>Anti-HIV in Human PBMCs</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>10% FBS</td>
<td>10% FBS + HSA + α1-AGP</td>
</tr>
<tr>
<td></td>
<td>EC_{50} (nM)</td>
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</tr>
<tr>
<td>GS-9224</td>
<td>1.3 ± 1.1</td>
<td>19 ± 5</td>
</tr>
<tr>
<td>GS-9160</td>
<td>1.2 ± 0.5</td>
<td>13 ± 6</td>
</tr>
<tr>
<td>RAL</td>
<td>7.5 ± 1</td>
<td>23 ± 5</td>
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a The EC_{50} for Control (10% FBS) and (10% FBS + HSA + α1-AGP) were run in parallel.
b Mean ± SD of at least 3 independent experiments.
c Mean ± SD of at least 3 donors.
d Mean ± SD of at least 2 independent experiments.
e The values of EC_{50} were calculated from EC_{50}.
Mean ± SE of at least 4 independent experiments using IN enzyme.

Table 3: Inhibition of Integrase Enzymatic Strand Transfer Activity and Integration in HIV-1 Infected SupT1 and MT-2 Cells.

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<tr>
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<tr>
<td>APV</td>
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Table 4: Resistance Profile of Mutant Viruses Conferring Resistance to INSTIs.

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Mean ± SE of at least 4 independent experiments using IN enzyme.

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</tr>
<tr>
<td>APV</td>
<td>1.5 ± 0.1</td>
<td>15</td>
</tr>
</tbody>
</table>

Table 4: Resistance Profile of Mutant Viruses Conferring Resistance to INSTIs.

Mean ± SE of at least 4 independent experiments using IN enzyme.

Table 3: Inhibition of Integrase Enzymatic Strand Transfer Activity and Integration in HIV-1 Infected SupT1 and MT-2 Cells.
a known natural polymorphism of IN that is not associated with resistance to INSTIs [43]. In fact, by passage 6, this mutation was no longer detectable. Since the level of resistance increased from 4.6-fold to >100-fold from passages 4 to 6 with the appearance of the additional mutation Q148K, we wished to determine whether combining IN mutations L74M, G140S and Q148K in a molecular clone of HIV-1 would recapitulate resistance to GS-9224. These mutations were introduced either individually or together into an infectious molecular clone of HIV-1. Interestingly, as individual mutations, Q148K conferred 42-fold resistance to GS-9224, L74M alone had no effect and G140S alone conferred 6-fold resistance (Table 6). When L74M and G140S were combined, the double mutant conferred 11-fold resistance, suggesting that L74M may potentiate resistance conferred by G140S, a property of L74M that we previously reported with regard to the E92V mutation [19]. Notably, the triple mutant L74M/G140S/Q148K conferred >100-fold resistance to GS-9224, a level consistent with that observed for passage 6 virus (Tables 5 and 6). All combinations of the tested mutations displayed comparable resistance levels to GS-9160 and GS-9224, but different levels of resistance against RAL. In particular, L74M did not potentiate G140S resistance against RAL. In addition, while the triple mutant conferred a higher level of resistance than Q148K alone against GS-9224 and GS-9160, it conferred a lower level of resistance than Q148K alone against RAL.

GS-9224 displays improved in vitro ADME properties compared to GS-9160

GS-9224 displayed a longer half-life than GS-9160 in human and dog hepatic microsomal extracts (Table 7). In addition, GS-9224 did not demonstrate any significant inhibition or induction of human CYP450 isoforms within clinically achievable range of concentrations (data not shown). When permeability was measured across a monolayer of Caco-2 cells, GS-9224 displayed higher permeability than GS-9160 in both apical to baso-lateral and baso-lateral to apical directions with a lower efflux ratio than GS-9160 or RAL (Table 7). Finally, the assessment of plasma protein binding by equilibrium dialysis against dog and human plasma showed that GS-9224 is more highly protein bound than GS-9160 both in dog plasma (~99.5% vs. ~93.5%) and in human plasma (~97.7% vs. ~89.6%) (Table 7).

Table 5: Resistance Phenotype of GS-9224 Selected Viral Pools.

<table>
<thead>
<tr>
<th>Compound</th>
<th>EC_{50} (nM)</th>
<th>CC_{50} (fold change relative to HIV-1 IIIb)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TFV</td>
<td>3.8 ± 0.8</td>
<td>3 ± 0.4 (0.9)</td>
</tr>
<tr>
<td>EFV</td>
<td>0.7 ± 0.1</td>
<td>1.3 ± 0.2 (1.9)</td>
</tr>
<tr>
<td>LPV</td>
<td>13.0 ± 4.0</td>
<td>27.0 ± 6.0 (2.1)</td>
</tr>
<tr>
<td>RAL</td>
<td>3.5 ± 0.6</td>
<td>11.0 ± 1.2 (3.1)</td>
</tr>
<tr>
<td>GS-9224</td>
<td>4.0 ± 2.0</td>
<td>18.6 ± 2.7 (4.6)</td>
</tr>
<tr>
<td>GS-9224</td>
<td></td>
<td>&gt;1000 (&gt;100)</td>
</tr>
</tbody>
</table>

* Mean ± SD of at least 6 independent experiments.

** Mean ± SD of at least 3 independent experiments.

The numbers in parentheses represent the fold-change over wild-type virus.
out to determine the specific uridine diphosphate transferase (UGT) isoforms responsible for the metabolism of GS-9224. The results showed that UGT 1A1 and 1A3 are the main UGT isoforms responsible for glucuronidation of both GS-9160 and GS-9224 (Table 8). The characterization of GS-9224 metabolites formed systemically both in vitro and in vivo supports the formation of a single phenolic glucuronic metabolite of GS-9224 (Figure 3).

Single-dose pharmacokinetics in healthy human subjects

After administration of single doses of GS-9224, 50 or 100 mg as an oral solution in the fasted state, maximum mean (CV%) concentrations of 110 ng/mL (51%) and 215 ng/mL (41%) were achieved at the 0.75 hour nominal time point, thereafter falling to <1.0 ng/mL in 8/8 of 11 subjects by the 12 hour time point for the 50 mg and 100 mg dose group, respectively. These data indicated rapid absorption (100%) and 7/8 (87.5%) subjects by the 12 hour time point for the 50 mg and 100 mg GS-9224 was indicative of the inhibition of proviral DNA integration. The elevation of 2-LTR circles in HIV-infected cells treated with GS-9224 was indicative of the inhibition of proviral DNA integration.

Discussion

GS-9224 is an analog of GS-9160, a previously reported investigational INSTI [19]. GS-9224 was designed in an effort to optimize the pharmacokinetic profile of GS-9160 while retaining its antiviral potency. The medicinal chemistry optimization focused primarily on the p-fluorobenzyl moiety of GS-9160. Profiling of various analogs led to the identification of GS-9224 that differs structurally from GS-9160 by the addition of one chloride atom and one fluorine atom on the p-fluorobenzyl moiety (Figure 1) [22].

Similar to GS-9160, GS-9224 exhibits potent and selective antiviral activity against HIV-1 and acts as an efficacious inhibitor of the IN strand transfer activity in an in vitro biochemical enzymatic assay. The elevation of 2-LTR circles in HIV-infected cells treated with GS-9224 was indicative of the inhibition of proviral DNA integration. In addition, GS-9224 profoundly reduced the number of integration events in treated cells as measured by Alu-LTR quantification of integration junctions without alteration of total viral DNA synthesis measured by the late-RT signal, demonstrating that GS-9224 is an authentic inhibitor of HIV-1 integration.

GS-9224 showed synergistic antiviral effect when combined with approved HIV drugs from different antiretroviral classes and retained antiviral potency against NRTI-, NNRTI- and PI-resistant HIV-1

<table>
<thead>
<tr>
<th>Compoundsa</th>
<th>EC50 (nM)</th>
<th>Fold-Change in EC50 Relative to HIV-1 IIIb</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HIV-1 IIb</td>
<td>L74M*</td>
</tr>
<tr>
<td>RAL</td>
<td>4.3</td>
<td>1.7</td>
</tr>
<tr>
<td>GS-9160</td>
<td>1.7</td>
<td>1</td>
</tr>
<tr>
<td>GS-9224</td>
<td>5.6</td>
<td>1</td>
</tr>
</tbody>
</table>

Table 6: Resistance Profile of Mutant Viruses Selected by GS-9224.

Table 7: In Vitro ADME Profiling of GS-9224 Relative to GS-9160 and RAL.

Table 8: Rate of Glucuronidation of GS-9224 by Major UGT Isoforms.

<table>
<thead>
<tr>
<th>UGT Isoform</th>
<th>Glucuronidation Rateb (pmol/min/mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1A1</td>
</tr>
<tr>
<td>GS-9160</td>
<td>16.7</td>
</tr>
<tr>
<td>GS-9224</td>
<td>75.4</td>
</tr>
</tbody>
</table>

a Selected by GS-9224 (this report), L-708,906 (32), S-1360 (33), L-870,810 (42) and GS-9160 (19).

b Selected by GS-9224 (this report) and L-Chicoric acid (29).

c Selected by GS-9224 (this report), S-1360 (31) and RAL (14, 39).

d The EC50 and fold-change represent the mean of at least two independent experiments performed in triplicate.

References

mutants, an observation consistent with its established mechanism of action. Characterizing the activity against a series of viruses expressing various INSTI resistance mutations revealed a similar resistance profile for GS-9224 and GS-9160. In comparison with RAL, both compounds showed lower degree of resistance against clinically relevant mutation Q148K, but higher level of resistance against E92Q and N155H mutants observed in patients treated with either RAL or EVG [44,45].

In vitro passaging of HIV-1 in the presence of gradually increasing concentrations of GS-9224 resulted in the selection of an IN mutant variant with >10-fold resistance to the selecting inhibitor and a similar cross-resistance to other INSTIs including RAL and the parent inhibitor GS-9160. Clonal sequencing of the resistant variant revealed the emergence of G140S and L74M mutations followed by the Q148K mutation. It is interesting to note that Q148K emerged as a primary mutation in patients treated with RAL who experienced virologic failure [3,4], but in the selection with GS-9224, Q148K emerged only after L74M and G140S. Thus, it appears that Q148K is a secondary mutation that augments resistance of L74M/G140S against GS-9224. In contrast, Q148K displays a higher degree of resistance to RAL as a primary single mutation compared to the triple mutant L74M/G140S/ Q148K selected by GS-9224.

The additional chlorine and fluorine atoms on the fluorobenzyl moiety of GS-9160 conferred to GS-9224 a moderately higher hepatic microsome stability, better absorption potential, lower efflux ratio and higher serum protein binding compared to GS-9160 (Table 7).

In healthy human subjects, GS-9224 rapidly achieved maximum plasma concentrations within one hour of administration as a solution in the fasted state. Thereafter, concentrations rapidly declined with a terminal elimination half-life of ~1.0 hour. These plasma concentrations within one hour of administration as a solution in the fasted state. Thereafter, concentrations rapidly declined with a terminal elimination half-life of ~1.0 hour. These human pharmacokinetic results were inconsistent with the target pharmacokinetic profile to provide sustained, therapeutic concentrations over a 24 hour dosing interval. As such, from a clinical pharmacology perspective, GS-9224 was deemed not suitable for QD dosing. In these clinically derived samples, the mean free fraction of GS-9224 was 7.5 to 10%, indicating lower in vivo protein binding than that estimated from in vitro plasma equilibrium dialysis for both dogs and humans. This could, at least in part, contribute to the fast systemic elimination of GS-9224 in humans. Single doses of GS-9224 at 50 mg and 100 mg were safe and well tolerated in healthy, adult, male and female subjects.

In conclusion, GS-9224 has retained the antiviral potency, mechanism of action and resistance profile of the predecessor compound, GS-9160. In addition, it exhibited improved in vitro ADME profile including higher metabolic stability, absorption potential and serum protein binding. However, these differential properties did not translate into an improved pharmacokinetic profile in healthy human volunteers. Results from the assessment of the single-dose pharmacokinetics revealed a profile of GS-9224 that would not be compatible with the once-daily oral dosing of GS-9224 due to faster than expected systemic elimination of the drug.

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References


