Title
Identification and Optimization of Thienopyridine Carboxamides as Inhibitors of HIV Regulatory Complexes

Permalink
https://escholarship.org/uc/item/9rz04807

Journal
Antimicrobial Agents and Chemotherapy, 61(7)

ISSN
0066-4804

Authors
Nakamura, Robert L
Burlingame, Mark A
Yang, Shumin
et al.

Publication Date
2017-07-01

DOI
10.1128/aac.02366-16

Peer reviewed
Identification and Optimization of Thienopyridine Carboxamides as Inhibitors of HIV Regulatory Complexes

Robert L. Nakamura,a,d Mark A. Burlingame,c Shumin Yang,b,d David C. Crosby,d Dale J. Talbot,a,d Kitty Chui,a Alan D. Frankel,d Adam R. Renslo

Advanced Genetic Systems, San Francisco, California, USAa; School of Medicine, Tsinghua University, Beijing, Chinaab; Small Molecule Discovery Center and Department of Pharmaceutical Chemistryc and Department of Biochemistry and Biophysics,d University of California, San Francisco, California, USA

ABSTRACT  Viral regulatory complexes perform critical functions during virus replication and are important targets for therapeutic intervention. In HIV, the Tat and Rev proteins form complexes with multiple viral and cellular factors to direct transcription and export of the viral RNA. These complexes are composed of many proteins and are dynamic, making them difficult to fully recapitulate in vitro. Therefore, we developed a cell-based reporter assay to monitor the assembly of viral complexes for inhibitor screening. We screened a small-molecule library and identified multiple hits that inhibit the activity of the viral complexes. A subsequent chemistry effort was focused on a thieno[2,3-b]pyridine scaffold, examples of which inhibited HIV replication and the emergence from viral latency. Notable aspects of the effort to determine the structure-activity relationship (SAR) include migration to the regiosomeric thieno[2,3-c]pyridine ring system and the identification of analogs with single-digit nanomolar activity in both reporter and HIV infectivity assays, an improvement of >100-fold in potency over the original hits. These results validate the screening strategy employed and reveal a promising lead series for the development of a new class of HIV therapeutics.

KEYWORDS antivirals, HIV, RNA, Rev

RNA-protein complexes are essential to the assembly and activity of many viral regulatory systems and represent an important target class for antiviral drug discovery. In HIV, the Rev-Rev response element (RRE) protein-RNA complex is one such target due to its essential activity in mediating the export of unspliced and partially spliced RNAs from the nucleus to the cytoplasm (1, 2). Disrupting the Rev-RRE interaction prevents the expression of late viral proteins and the packaging of viral RNA, thus inhibiting virus replication.

Rev is a 116-amino-acid RNA-binding protein that is expressed from fully spliced mRNAs early in the virus life cycle (3). Rev binds the RRE, a highly structured ~350-nucleotide (nt) RNA element encoded within the env gene (reviewed in references 2 and 4). Current models suggest that about six Rev molecules bind the RNA in order to properly position two of the nuclear export sequences on Rev for binding to a dimer of the Crm1-RanGTP export complex (5, 6). This complex is then exported through the nuclear pore, after which it disassembles in the cytoplasm to allow translation of the late HIV proteins and packaging of the viral genome (2, 7).

The formation of the export complex is driven by several critical intermolecular interactions. Rev binds the RNA primarily through its arginine-rich motif (ARM), an α-helical domain that forms several important hydrogen bonds with the RNA (8, 9). The nuclear magnetic resonance (NMR) structures of the Rev peptide complexed to RRE IIB or to an RNA aptamer and a recent cocrystal structure of a Rev-RRE dimer (10) show that...
the α-helix binds in the widened major groove of the high-affinity IIB site, primarily using arginine and asparagine side chains for base-specific recognition (11–13). The cocrystal structure also shows that a glutamine at position 51 interacts symmetrically across the dimer interface to help stabilize the complex (10). Flanking the ARM of Rev are hydrophobic oligomerization domains that drive the formation of a stable Rev dimer, and Rev dimers in turn form higher-order structures through hydrophobic multimerization domains that are located opposite the oligomerization domain, ultimately assembling a very high affinity (picomolar) complex (14–16). Given the essential nature of this assembly and the requirement for high-affinity binding, small molecules that interfere with any of these interfaces are expected to block formation of competent export complexes and to be of potential therapeutic benefit.

To identify small-molecule inhibitors of the Rev-RRE complex, we developed a high-throughput assay based upon the established Tat hybrid assay platform for characterizing RNA-protein or RNA-peptide interactions in mammalian cells (17, 18). This system takes advantage of the function of the HIV transcription activator Tat, which binds to the TAR RNA hairpin located at the 5′ end of the viral transcripts (19, 20) and enhances the processivity of RNA polymerase II transcription complexes initiated at the HIV long terminal repeat (LTR) (21–23). The Tat activation domain can be used to activate reporter expression via heterologous RNA-protein interactions (24, 25) (Fig. 1A). Reporter activity is proportional to RNA-binding affinity, and thus the assay has been used extensively to characterize RNA-protein interactions, including HIV Tat-TAR, bovine immunodeficiency virus (BIV) Tat-TAR, and HIV Rev-RRE, and to screen for novel proteins that bind the Mason-Pfizer monkey virus (MPMV) constitutive transport element (CTE) (17, 26–32).

We engineered the Tat hybrid platform with the Rev-RRE interaction and used it to screen a targeted small-molecule compound library. We hypothesized that compounds

![Screening strategy to identify inhibitors of RNA-protein complexes.](image-url)
containing a carboxamide moiety might mimic the essential asparagine or glutamine side chains and compete for Rev-Rev dimerization or Rev binding to the RRE RNA, and we thus constructed a library of ~4,500 carboxamide-containing small molecules selected from a larger-diversity library. Here, we describe the discovery and structure-activity studies of a thienopyridine inhibitor scaffold in which an unsubstituted carboxamide function was found to be essential for activity. Optimized thienopyridine analogs exhibited low-nanomolar potencies in multiple reporter-based assays as well as in HIV replication assays. Interestingly, related thienopyridine analogs were identified independently as Rev inhibitors using an orthogonal screening approach (33).

RESULTS

We developed a Tat hybrid-based screening assay to identify small molecules that target RNA-protein interactions in general and the Rev-RRE interaction in particular (Fig. 1B). To screen small molecules, we engineered a stable Tat hybrid reporter cell line encoding the RRE IIB RNA target and driving a firefly luciferase (FFL) reporter. The reporter exhibited a good signal-to-noise ratio and was compatible with multiwell plates, robotic liquid handling, and standard luminescence plate readers. The Tat-Rev expression plasmid was then integrated into the reporter cell line, and clonal cell lines were selected that constitutively expressed luciferase to high levels, indicative of delivery of Tat via the heterologous RNA-protein interaction. We validated the assay using small interfering RNA (siRNA) knockdowns, targeting the Tat activation domain, and with 3,6-diaminoacridine, a known small-molecule inhibitor of the Rev-RRE interaction that served as a positive control (see Fig. S1 in the supplemental material) (34).

The RRE IIB-Rev screening cell line was plated in 384-well assay plates, and compounds were added to the cells to a final concentration of 30 μM. Compounds were screened in duplicate, and the luciferase activity of each compound was highly consistent (typically <10% deviation in activity) (Fig. S2). In parallel, MTT [3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyl-2H-tetrazolium bromide] cell viability assays were performed to assess the toxicity of compounds and minimize the level of false positives. Screening data for the initial library screen is shown in Fig. 1C.

From the primary screen, we obtained 11 validated hits with confirmed dose-dependent activity and a lack of toxicity in the MTT assay. Of these 11 compounds, two were benzopyrans, one was a tetrasubstituted thiophene, and the remaining eight hits possessed a common thieno[2,3-b]pyridine core (Fig. S3). To further validate the hits, we used an electrophoretic mobility shift assay (EMSA) to evaluate which of the three hit scaffolds were capable of disrupting the Rev-RRE IIB interaction in vitro. Only the thiophene scaffold disrupted the complex, as determined by EMSA (Fig. S4). This result demonstrated that the Tat hybrid screening method could identify bona fide inhibitors of RNA-protein interactions although our experience with EMSA raised questions about this method. We observed that compounds that disrupted the RNA-protein complex by EMSA, namely, 3,6-diaminoacridine, neomycin B, and thiophenes, were toxic to cells. These types of relatively nonspecific intercalating agents (e.g., acridines) and RNA-binding compounds (e.g., aminoglycosides) can be readily picked up as binding inhibitors in EMSAs but may be missed in cell-based assays because of their high toxicity. Conversely, the lack of EMSA inhibition by other types of compounds that scored well in the cell-based assay does not necessarily rule out Rev-RRE as their target since the in vitro EMSA lacks other proteins required for Rev function (e.g., Crm1 and Ran) and potential accessory factors (e.g., hnRNPs and chaperones) that may alter the nature of Rev-RRE complexes for inhibitors to act on. Other experimental evidence supporting Rev-RRE as a plausible target of thienopyridines is provided below. Because of a superior toxicity profile and a nascent structure-activity relationship (SAR) profile, we focused our efforts on the thienopyridine scaffold.

Compounds 1a to c (Fig. 2) are representative of the original thienopyridine hits, which varied in terms of substitution on the pyridine ring but universally possessed amino and carboxamide substitution on the thiophene ring. Commercially available analogs of compound 1a bearing a carboxylic acid (1d) or methyl ketone (1e) in place...
of the carboxamide were purchased and found not to be significantly active in the Tat hybrid assay. This suggested the importance of the carboxamide function for activity in the Tat hybrid reporter cell line (Fig. 2 and 3; Table 1). Evaluation of additional commercial analogs suggested that modification of the 4 or 5 position of the thien-
opyridine ring was generally well tolerated, and this afforded some confidence that improvements in potency could be realized with the synthesis of additional analogs (Table S1).

Because of the reasonable activity of thienopyridine compounds in the Tat hybrid reporter assay, low toxicity in the MTT assay, and a nascent SAR profile, we embarked on structure-activity studies of the thienopyridine series. One of the first analogs synthesized was compound 2a derived from a thieno[2,3-c]pyridine ring system that is regioisomeric with the original hit compounds (Fig. 2). Gratifyingly, 2a exhibited submicromolar potency in both the Tat hybrid assay and in a viral replication assay in U1 cells (Table 1). The U1 cells contain an integrated and inducible HIV-1 provirus, meaning that inhibition of viral replication in these cells is indicative of compound action subsequent to retroviral genome integration, as expected for a Rev-RRE inhibitor. As the potency of analog 2a was similar to that of its directly analogous thieno[2,3-b]thionopyridine congener 1f in both assays (Fig. 2 and Table 1), we focused our SAR studies on the comparatively unexplored thieno[2,3-c]thionopyridine scaffold represented by compound 2a.

All thieno[2,3-c]pyridine analogs described herein were prepared by one of the three synthetic approaches described below (Fig. 4) (35). Thus, the preparation of the group 2 amine-bearing analogs involved SNAr reaction of secondary amines with 3,5-dichloro-4-pyridinecarbonitrile, followed by reaction with 2-mercaptoacetamide in the presence of sodium methoxide. Aryl and heteroaryl-substituted group 3 analogs were prepared similarly, but beginning with Suzuki coupling reactions. Finally, des-amino group 4 variants were prepared from 3,5-dihalopyridinecarboxaldehyde via initial formation of the thieno[2,3-c]pyridine ring as before, followed by Suzuki coupling (36).

New analogs were evaluated in both the Tat hybrid assay, which reports on the Rev-RRE interaction, and in the viral activation assay in U1 cells (Table 1). Representative dose-response curves in the U1 assay are shown for analogs 1a, 2b, and 4e (Fig. 5 and Table S2). We found that the activities of ~50 synthetic thienopyridine analogs were well correlated in the Tat hybrid and U1 assays (Fig. 6), suggesting that compound action remained on target during optimization and consistent with the postintegration pharmacology expected of a Rev-RRE inhibitor.

To further explore amine substitution as in analog 2a, additional analogs such as the piperidine 2b, morpholine 2d, and N-Me piperazine analog 2e were prepared and evaluated (Fig. 7 and Table 1). Piperidine analog 2b was nearly 10-fold more potent than the dimethylamino analog 2a, an effect that appears to derive from the aliphatic

| Table 1 Activity of thienopyridine group 1 and 2 analogs in reporter and antiviral assays |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Compound | Tat hybrid assay EC50 (nM) | U1 replication assay IC50 (nM) | Rev reporter assay EC50 (nM) | HIV-1 replication assay IC50 (nM) | LipE \(\text{a}^{\text{b}}\) |
| 1a | 2,170 | 1,473 | 4.7 |
| 1b | 1,160 | 528 | 5.2 |
| 1c | 700 | 750 | 4.5 |
| 1d | >100,000 | | |
| 1e | 31,200 | | 2.4 |
| 1f | 210 | 659 | 530 | 8.2 |
| 2a | 710 | 450 | 5.8 |
| 2b | 65 | 28 | 680 | 77 | 5.9 |
| 2c | 330 | 391 | 6.5 |
| 2d | 930 | | 6.2 |
| 2e | 7,620 | | 5.3 |
| 2f | 140 | 219 | 634 | 5.2 |
| 2g | 370 | 173 | 5.75 |
| 2h | 350 | 219 | 5.0 |
| 2i | 70 | 13 | 5.3 |
| 2j | 100 | 34 | 4.6 |
| 2k | 730 | 1,087 | 4.8 |

\(\text{a}^{\text{b}}\) LipE = \(\text{p}K_I - \text{clogP}\); \(\text{p}K_I\) was estimated from the Tat hybrid assay EC50; clogP values were determined in Vortex (Dotmatics) using the calculated property XlogP.
nature of the piperidine ring, substitution of which (2c) or replacement with a morpholine (2d) or piperazine (2e) ring resulted in less potent analogs. Spirocyclic (2g) and geminal-difluoro (2h) substitutions at the 4 position proved inferior to results with the parent piperidine 2b. Tetrahydroquinoline (2i) and phenyl ether (2j) analogs were equipotent to 2b but with their additional mass and lipophilicity produced inferior lipophilic efficiency (\(\text{LipE} = p\_K_i - \text{clogP}\)) (37). Finally, introduction of a heteroaryl ring at the 3 position of the piperidine ring as in compound 2k was poorly tolerated. Although N-acylpiperazine analog 2f was notably more potent than the basic N-methylpiperazine analog 2e, the exploration of additional amide variants did not yield useful gains in potency.

Having identified analogs like 2b and 2l that were up to 10-fold more potent than 2a, we next explored 4-aryl-substituted thienopyridine analogs either with (3a to j) or without (4a to j) a 3-amino group (Fig. 8 and Table 2). In the case of substituted phenyl rings, we found substitution at the para and/or meta position(s) was favored over ortho substitution (compare compounds 4c and 4d). Moderately electron-rich analogs like

---

**Fig 4** Synthesis of thieno[2,3-c]pyridine analogs 2 to 4. Conditions: a, 1 to 2 equivalents R1R2NH, Et3N, dimethylformamide, 25 to 40°C, 12 to 24 h; b, 2 equivalents 2-mercaptoacetamide, 10 wt% in MeOH/NH₃, 2 equivalents NaOMe, dimethylformamide, \(\mu\text{W}, 80°C, 10\) to 60 min; c, 1 to 2 equivalents Ar-B(OH)₂, 0.025 equivalent PCy₃, 0.01 equivalent Pd₂(dba)₃, 1.7 equivalents K₂CO₃, dioxane, \(\mu\text{W}, 150°C, 30\) min; d, 0.75 equivalent 2-mercaptoacetamide, 10 wt% in MeOH/NH₃, 1.2 equivalents NaOMe, dimethylformamide, \(\mu\text{W}, 80°C, 10\) to 60 min.

**Fig 5** Activity of thienopyridine analogs 1a, 2b, and 4e in the U1 activation assay. U1 cells were plated in 96-well plates and activated by the addition of phytohemagglutinin. The test compounds were added to the cells at doses ranging from 1 nM to 3,160 nM in triplicate, and supernatants were collected 72 h later. p24 values were determined by ELISA, and average values are shown.
3a/4a and 3e/4e were more potent than either electron-deficient congeners (4c) or more electron-rich analogs like 3f/4f. Consistent with substitution effects in the piperazine series (2k), bulky substitution at the 3 position (meta) as in 3j/4j was poorly tolerated. The most consistent effect in this series was the favorable effect of removing the 3-amino group (as in group 4 analogs, X represents H). Particularly potent were des-amino analogs bearing a moderately electron-rich aryl ring, as in the analogs 4a and 4e, which exhibited low-nanomolar potency in both the Tat hybrid and U1 assays and were the most potent analogs evaluated (Table 2). Analogs 4a and 4e combine exceptional potencies with reasonable calculated lipophilicities, thus affording LipE values of 5.0 and 5.3, respectively. These analogs, along with the potent but more lipophilic analogs 4g and 4h, were evaluated in an HIV-1 replication assay. Jurkat cells were infected with HIV-1 laboratory isolate NL4-3 in the presence of the test compounds, and viral production was monitored by collecting viral supernatants and measuring p24 levels by enzyme-linked immunosorbent assay (ELISA) (Fig. 9 and Table S3). All four analogs displayed low-nanomolar efficacy in this assay, and moreover, their rank order potencies in the infectivity assay were correlated with 50% inhibitory concentrations (IC50s) in the U1 activation assay (Table 2). Replication curves for the des-amino analogs 4e and 4h are shown in Fig. 9B and C. These compounds exhibited IC50s of 3.9 nM and 3.4 nM, respectively, in the HIV-1 replication assay.
assay compared to 50% toxic concentration (TC_{50}) values of 19,200 and 20,500 nM in the MTT cell viability assay. For comparison, AZT exhibited an IC_{50} of 3 nM and TC_{50} of >100,000 nM in these same assays. The piperazine analog 2b was also evaluated in the HIV-1 replication assay and exhibited activities in the mid- to high-nanomolar range (77 nM) and modest toxicity (41,400 nM) (Fig. 9A).

We next tested compound 4e in replication assays using primary cells and clinical isolates of HIV-1. Peripheral blood mononuclear cells (PBMCs) were stimulated with phytohemagglutinin (PHA) and infected with several clinical isolates of HIV-1 in the presence of the compounds. IC_{50}s for the compound were determined by collecting viral supernatants and measuring p24 levels by ELISA (Table 3). Results of this experiment are shown in Fig. 10 where the dose-response and toxicity curves for isolate 93BR021 (CCR5-tropic, group M, subtype B) are shown. Here, we observed an IC_{50} of 55 nM and TC_{50} of 32,300 nM. AZT was again used as a positive control and exhibited an IC_{50} of 18.5 nM, and the TC_{50} was determined to be >50,000 nM by MTT assay.

We next evaluated analogs 2b, 2f, 4a, and 4e in a panel of standard in vitro absorption, distribution, metabolism, and excretion (ADME) assays to assess drug-like properties and the potential for efficacy in animals. All four analogs are rule-of-five compliant and exhibited moderate to high permeability across Caco-2 monolayers, a

![Structures of thieno[2,3-c]pyridine analogs of groups 3 and 4.](image)

**TABLE 2** Activity of thienopyridine group 3 and 4 analogs in reporter and antiviral assays

<table>
<thead>
<tr>
<th>Compound</th>
<th>Tat hybrid assay EC_{50} (nM)</th>
<th>U1 replication assay IC_{50} (nM)</th>
<th>Rev reporter assay EC_{50} (nM)</th>
<th>HIV-1 replication assay IC_{50} (nM)</th>
<th>LipEx</th>
</tr>
</thead>
<tbody>
<tr>
<td>3a</td>
<td>70</td>
<td>57</td>
<td>4.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3b</td>
<td>140</td>
<td>38</td>
<td>4.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3e</td>
<td>200</td>
<td>91</td>
<td>4.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3f</td>
<td>330</td>
<td>277</td>
<td>4.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3j</td>
<td>4,700</td>
<td></td>
<td>1.15</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4a</td>
<td>10</td>
<td>11</td>
<td>4.5</td>
<td></td>
<td>5.0</td>
</tr>
<tr>
<td>4c</td>
<td>140</td>
<td>108</td>
<td>4.0</td>
<td></td>
<td>4.0</td>
</tr>
<tr>
<td>4d</td>
<td>560</td>
<td>946</td>
<td>3.5</td>
<td></td>
<td>3.5</td>
</tr>
<tr>
<td>4e</td>
<td>8</td>
<td>9</td>
<td>3.9</td>
<td></td>
<td>5.3</td>
</tr>
<tr>
<td>4f</td>
<td>70</td>
<td>65</td>
<td>4.5</td>
<td></td>
<td>4.5</td>
</tr>
<tr>
<td>4g</td>
<td>41</td>
<td>66</td>
<td>6.4</td>
<td></td>
<td>4.2</td>
</tr>
<tr>
<td>4h</td>
<td>34</td>
<td>8.9</td>
<td>3.4</td>
<td></td>
<td>3.2</td>
</tr>
<tr>
<td>4i</td>
<td>160</td>
<td>7.7</td>
<td>2.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4j</td>
<td>150</td>
<td>368</td>
<td>1.9</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*LipE* = pK_{a} – clogP, pK_{a}, was estimated from the Tat hybrid EC_{50}, clogP values were determined in Vortex using the calculated property XlogP.
model of intestinal absorption (Table 4). Analogs 4a and 4e, with only two hydrogen bond donors, were more permeable but also notably less soluble than analogs 2b and 2f. The more lipophilic analogs 4a and 4e also appear more prone to oxidative metabolism in the presence of human liver microsomes. This is perhaps unsurprising given the presence of methoxyphenyl (4a) and benzylic methylene (4e) functions, both potential sites of oxidative metabolism.

**FIG 9** Replication assays with HIV-1 isolate NL4-3 in Jurkat cells. Replication spreading assays were performed by infecting Jurkat cells with NL4-3 in the presence of compound at doses ranging from 3.16 nM to 1,000 nM in triplicate. Supernatants were collected every 48 to 72 h, and p24 values were determined by ELISA. Average replication curves are shown for the indicated compounds.
DISCUSSION

Here, we used a screening approach based on the HIV LTR Tat hybrid system to identify small molecules putatively targeting a protein-RNA interaction. A focused library of carboxamide-containing compounds was screened based on the expectation that such molecules can interact with RNA bases and could potentially disrupt or alter Rev-RNA binding interactions. Of the various hits identified in the screen, a series of thienopyridine carboxamides emerged as the most promising hit scaffold. We then used an HIV reporter cell line and the U1 replication assay to validate the screening hits and to evaluate an additional ~100 commercial analogs. These analyses revealed that an unsubstituted carboxamide group at the 2 position of the thienopyridine ring was essential for activity. Furthermore, substitution on the thienopyridine ring was well tolerated, particularly at positions 4 and 5. This initial survey of commercially available analogs provided a preliminary SAR profile that encouraged further synthetic efforts on the scaffold.

A variety of analogs was prepared based on a regioisomeric thieno[2,3-c]pyridine ring system. Substitution of the 4 position with N-linked heteroaliphatic rings (group 2 compounds) or C-linked aryl and heteroaryl rings (groups 3 and 4) was well tolerated, with the C-linked aryl analogs in general possessing superior potencies. Most significantly, we found that the 3-amino group is not essential for activity and that, in fact, des-amino analogs possess notably superior potencies in both the Tat hybrid reporter assay and the U1 assay. The elimination of two hydrogen bond donors in such analogs likely improves intrinsic cell permeability, which may wholly or partly explain the enhanced potency of group 4 analogs in cell-based assays. Of the des-amino analogs, compound 4e showed the best potencies and therapeutic indices in the reporter and U1 assays and in the HIV replication assays, with a 50% effective concentration (EC_{50})

<table>
<thead>
<tr>
<th>Viral isolate</th>
<th>Coreceptor(s)</th>
<th>Compound 4e IC_{50} (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ba-L</td>
<td>CCR5</td>
<td>48.8</td>
</tr>
<tr>
<td>93BR021</td>
<td>CCR5</td>
<td>54.8</td>
</tr>
<tr>
<td>93BR028</td>
<td>CCR5</td>
<td>9.2</td>
</tr>
<tr>
<td>92TH014</td>
<td>CCR5</td>
<td>19.6</td>
</tr>
<tr>
<td>92UG005</td>
<td>CXCR4</td>
<td>70.8</td>
</tr>
<tr>
<td>92HT599</td>
<td>CXCR4</td>
<td>25.9</td>
</tr>
<tr>
<td>LAI</td>
<td>CXCR4</td>
<td>291</td>
</tr>
<tr>
<td>92HT593</td>
<td>CXCR4, CCR5</td>
<td>13.4</td>
</tr>
</tbody>
</table>

**TABLE 3** Activity of thienopyridine analog 4e in replication assays in PBMCs

**FIG 10** Replication assay with HIV-1 primary isolate 93BR021 in PBMCs. Replication assays were performed in PBMCs by stimulating cells with phytohemagglutinin and then infecting them with clinical isolates of HIV. Compound 4e was added to PBMCs at doses ranging from 1 nM to 1,000 nM in triplicate or more, and supernatants were collected every 48 to 72 h. p24 values were determined by ELISA. Average replication curves are shown in blue, and toxicity is shown in green.
of 8 nM in the reporter assay and IC_{50}s of 9 nM and 3.9 nM in the U1 and HIV replication assays, respectively (Table 2). By comparison, the most potent of the N-linked analogs, piperazine 2b, displayed an EC_{50} of 65 nM in the reporter assay and IC_{50}s of 28 nM and 77 nM in the U1 and HIV replication assays, respectively (Table 1). Overall, our structure activity studies improved potencies from the low-micromolar to the low-nanomolar regime, likely by improved target-level binding affinity and improved cellular permeability. While further optimization of in vivo drug-like properties will be required, leads like analog 4e appear promising as a starting point for such efforts.

Thienopyridine analogs that are closely related to our initial screening hits were described as inhibitors of Rev-RRE by another group of investigators (33) using a reporter assay that is dependent upon the activity of Rev (38). We therefore tested selected compounds from our SAR studies (1f, 2b, 4a and 4e) in a similar Rev-dependent reporter assay (Tables 1 and 2). Several additional experiments were described that strongly suggest that Rev is the target of the thienopyridine analogs, including additional reporter experiments that showed inhibition of a Rev-dependent, but not a CTE-dependent, reporter as well as experiments that showed that the protein composition of inhibitor-treated samples was consistent with an inhibitor of Rev function (33).

Biochemical assays including EMSAs did not show inhibition of the Rev-RRE interaction by the thienopyridine class in vitro. These assays generally capture a snapshot of Rev activity and lack critical interactions with cellular cofactors such as Crm1 and Ran. We have found that compounds that inhibit formation of RNA-protein complexes are often toxic, suggesting that EMSAs or other in vitro assays may lack the sensitivity to reveal subtle effects of small-molecule inhibitors. In some cases, such as with 3,6-diaminoacridine and neomycin B, inhibition of Rev-RRE formation is observed by EMSA, but both are toxic at therapeutic concentrations against HIV. Similarly, the thiophene class identified by the small-molecule screen also inhibits the Rev-RRE interaction, as determined by EMSA (see Fig. S4 in the supplemental material), but several of the commercially available thiophene analogs showed substantial toxicity and thus were not pursued further. Interestingly, SAR analysis of the thiophenes and thienopyridines identified the thienopyridine as a better scaffold that likely maintained similar binding characteristics (39).

Although our studies and the previous work implicate Rev as the likely target of thienopyridine carboxamides, other targets cannot be ruled out. For example, our screen is dependent upon the activity of the HIV LTR, and thus an inhibitor of HIV transcription would also produce a positive response in the assay. However, this would appear unlikely because we showed, using the Tat hybrid assay (Fig. S5), that the thienopyridine compound 4a specifically targets the Rev-RRE reporter but not the HIV Tat-TAR reporter. The previous screen did not use the HIV LTR, and these investigators further uncovered a thienopyridine resistance mutation in the RRE, consistent with the Rev-RRE interaction as the target of thienopyridines (40). We have also generated this RRE-defective virus and shown that it is resistant to compound 4e (Fig. S6). Though strongly supportive that Rev is the target of the compounds, these data do not explicitly rule out the possibility of alternative targets; for example, thienopyridines may directly target structured RNA, or multiple regions of the HIV genome may be targeted by the compounds in addition to the RRE. In summary, we identified and optimized a

### TABLE 4 In vitro ADME properties of selected thienopyridine analogs

<table>
<thead>
<tr>
<th>Compound</th>
<th>Tat hybrid assay EC_{50} (nM)</th>
<th>Aqueous solubility at pH 7.4 (nM)</th>
<th>P_{app} of Caco-2 cells (nm/s)</th>
<th>HLM t_{1/2} (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2b</td>
<td>65</td>
<td>34,000</td>
<td>505</td>
<td>28</td>
</tr>
<tr>
<td>2f</td>
<td>140</td>
<td>258,000</td>
<td>97</td>
<td>&gt;60</td>
</tr>
<tr>
<td>4a</td>
<td>10</td>
<td>440</td>
<td>594</td>
<td>8</td>
</tr>
<tr>
<td>4e</td>
<td>8</td>
<td>280</td>
<td>666</td>
<td>18</td>
</tr>
</tbody>
</table>

*aP_{app}, apparent permeability.

bHLM t_{1/2}, half-life in human liver microsomes.
class of thienopyridine carboxamides that potently inhibits HIV replication in cells. This work represents an early but important step toward the development of a new class of HIV therapeutics that could eventually find a place in future HIV combination therapies. While the weight of evidence suggests that these compounds act by inhibition of the Rev-RRE interaction, further studies will be required to definitively establish their molecular pharmacology.

MATERIALS AND METHODS

**Small-molecule screen.** RNA reporter plasmids were constructed using the pHIV LTR vector (18). To express HIV Tat fusion proteins, we constructed a mammalian codon-optimized pcDNA-Tat vector containing the first 48 amino acids of HIV Tat with a fusion to HIV Rev (residues 3 to 70). HIV Tat fusion proteins have been previously described (17, 41). A HeLa cell line expressing the pcDNA3 HIV-1 LTR-RRE III-FFL reporter was generated to obtain a consistent background for library screens. The plasmid was transfected into HeLa cells, and stable integrants were selected using neomycin (G418) (800 μg/ml) for 10 days. Resistant cells were transfected with the pcDNA3 Tat-Rev3-70 plasmid harboring a hygro- mycin resistance marker. Individual clonal cell lines were assayed for luciferase signal, and 3,6- diaminoacridine, a nonspecific small-molecule inhibitor of the reporter, was used to identify cell lines with high signal-to-noise ratios (34). A Z’ value of 0.75 was obtained for the cell line used for the small-molecule screen (see Fig. S1 in the supplemental material).

A library of approximately 4,500 carboxamide-containing compounds was identified and purchased from ChemDiv. Approximately 5,000 cells were plated in white 384-well plate, and compounds were added to the cells to a final concentration of 30 μM using a Biomek FX robotic liquid handler (Beckman Coulter). The cells were incubated with test compounds for 48 h and assayed for luciferase activity (Bright Glo; Promega). The cytotoxicity of each compound was determined in parallel via 3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyl-2H-tetrazolium bromide (MTT)-based cell viability assays (42). Additional compounds structurally related to screening hits were purchased from Hit2Lead/Chembridge, Asinex, and Specs.

The test compounds that displayed activity were reconfirmed in duplicate or triplicate at multiple concentrations using the Tat hybrid reporter cell lines or by transient transfection. Previously characterized thienopyridines and 3,6-diaminoacridine (34) were used as controls. Cells were counted and plated into white 384-well or 96-well cell culture-treated assay plates, and compounds were added. After 48 h, the cells were prepared for luminescence assays that were performed on a plate reader (Tecan Evolution or MD Analyst). The cytotoxicity of each compound was determined in parallel via MTT-based cell viability assays.

**Rev reporter assay.** 293T cells were plated in 96-well plates and cotransfected with an HIV RRE (Rev-dependent) reporter plasmid (43) and a pSV2 Rev expression plasmid. Compounds were incubated with the cells for 48 h, and reporter activity was assessed by p24 ELISA. Leptomycin B, an inhibitor of the Crm1 pathway that inhibits Rev-dependent export, was included as a positive control. The cytotoxicity of each compound was determined in parallel via MTT-based cell viability assays.

**U1 activation assay.** U1 cells obtained from the AIDS Research and Reference Reagent Program were maintained under standard culture conditions in RPMI 1640 medium supplemented with 10% fetal bovine serum (heat inactivated), 2 mM l-glutamine, 100 U/ml penicillin, and 100 μg/ml streptomycin. U1 cells contain an integrated copy of a phorbol ester-inducible HIV-1 provirus, and addition of phytotermagglutinin (PHA) is used to induce virus production. On the day of the assay, the U1 cells were activated with PHA, and 2.5 × 10^5 cells were plated in 96-well plates. Test compounds diluted in medium were immediately applied to the cells at final concentrations of from 1 to 10,000 nM. Cultures were incubated for 3 days, and supernatants were harvested. Cell-free virus production was measured by p24 ELISA in culture supernatants. Compound toxicity was determined by MTT cell viability assay in parallel assays.

**Virus spreading assay.** Jurkat (E6-1) cells (NIH AIDS Reagent Program) were cultured in RPMI medium supplemented with 25 mM HEPES, pH 7.4, 10% heat-inactivated fetal calf serum (HyClone, Waltham MA), and 1% penicillin-streptomycin. A total of 2.5 × 10^6 Jurkat cells were inoculated with 250 pg of p24 in 250 μl of medium in a 96-well flat-bottom polystyrene cell culture microplate. Input virus was removed after 18 h via washing cells in phosphate-buffered saline (PBS). Cells were resuspended in 250 μl of medium containing compounds at the concentrations described in Fig. 9. Cells were incubated at 37°C in 5% CO2, supernatant samples were removed every 48 h, and medium was replaced with fresh medium containing the compounds. Viral replication was determined using an ELISA to quantify p24 viral capsid protein in the culture supernatant. Mouse monoclonal and rabbit polyclonal anti-p24 antibodies used in the ELISA were obtained from the NIH AIDS Reagent Program. Toxicity was monitored using the MTT cell viability assay.

**HIV-1 replication assay in PBMCs.** Virus isolates were obtained from the NIH AIDS Reagent Program. Human PBMCs, seronegative for HIV and hepatitis B virus (Astarte Biologics, LLC), were stimulated in R-3 medium (RPMI 1640 medium with 25 mM HEPES and l-glutamine, 20% heat-inactivated fetal bovine serum, 1% penicillin-streptomycin, 20 IU/ml interleukin-2 [IL-2]) with 5 μg/ml phytohemagglutinin (PHA) for 48 to 72 h. Stimulated PBMCs were resuspended and diluted in fresh R-3 medium and added to 96-well plates at 5 × 10^4 cells/well. The 50% tissue culture infective dose (TCID50) of each virus stock was measured by endpoint dilution assay as described by Reed and Muench (44). Spinoculation (1,200 × g, 2 h) was applied to improve the efficiency of infection.
Cells were infected with 40 TCID$_{50}$ of virus stock under spinoculation in the presence of different concentrations of test compounds (biological quintuplicate wells/concentration) incubated at 37°C in 5% CO$_2$ for 7 days. On day 4, half of the supernatant was removed and replaced with fresh R-3 medium containing the appropriate concentration of test compounds. On day 7, cell-free supernatant samples were collected for analysis of p24 antigen expression measured by ELISA as described above. Antiviral activity was assessed by the inhibition of p24 expression.

In vitro ADME assays. Aqueous solubility studies at pH 7.4 in phosphate buffer, Caco-2 permeability studies, and human liver microsome (HLM) stability studies were performed by Absorption Systems (Exton, PA) using their standard methods.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/AAC.02366-16.

SUPPLEMENTAL FILE 1, PDF file, 2.7 MB.

ACKNOWLEDGMENTS

We thank David Rekosh and Marie-Louise Hammarskjold of the University of Virginia for providing the Rev reporter construct and control small molecules, Kip Guy for critical help in designing the library screen, and members of the Frankel Lab for providing critical feedback and comments.

This work was supported by National Institutes of Health grants CA103407 to A.D.F. and R.L.N., AI076143 to R.L.N. and A.D.F., AI076087 to A.R.R. and R.L.N., and P50GM082250 to A.D.F., by a U.S. Treasury Department Qualified Therapeutic Discovery Grant to Advanced Genetics Systems, and by China Scholarship Council financial support to S.Y.

REFERENCES


