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Identification and Characterization of Small Molecules That Inhibit Intracellular Toxin Transport

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Shiga toxin (Stx), cholera toxin (Ctx), and the plant toxin ricin are among several toxins that reach their intracellular destinations via a complex route. Following endocytosis, these toxins travel in a retrograde direction through the endosomal system to the trans-Golgi network, Golgi apparatus, and endoplasmic reticulum (ER). There the toxins are transported across the ER membrane to the cytosol, where they carry out their toxic effects. Transport via the ER from the cell surface to the cytosol is apparently unique to pathogenic toxins, raising the possibility that various stages in the transport pathway can be therapeutically targeted. We have applied a luciferase-based high-throughput screen to a chemical library of small-molecule compounds in order to identify inhibitors of Stx. We report two novel compounds that protect against Stx and ricin inhibition of protein synthesis, and we demonstrate that these compounds reversibly inhibit bacterial transport at various stages in the endocytic pathway. One compound (compound 75) inhibited transport at an early stage of Stx and Ctx transport and also provided protection against diphtheria toxin, which enters the cytosol from early endosomes. In contrast, compound 134 inhibited transport from recycling endosomes through the Golgi apparatus and protected only against toxins that access the ER. Small-molecule compounds such as these will provide insight into the mechanism of toxin transport and lead to the identification of compounds with therapeutic potential against toxins routed through the ER.

Bacterial and plant toxins are significant agents of human disease and potential vehicles for bioterrorism. Though their intracellular targets are diverse, a common and essential step in their virulence is the ability to reach the cytosol, where most toxins exert their enzymatic effects. The bacterial exotoxins Shiga toxin (Stx) and cholera toxin (Ctx), as well as the plant toxin ricin, have drawn particular interest for their unique retrograde transport following endocytosis. Members of the AB toxin group, these toxins consist of a receptor-binding B subunit and an enzymatic A subunit. In contrast to anthrax and diphtheria toxins, AB toxins that enter the cytosol directly from early endosomes in a pH-dependent manner (18, 28, 35), these membrane-bound toxins bypass the late endocytic pathway by retrograde transport from early or sorting endosomes to the trans-Golgi network (TGN) (14, 15, 37, 60). From the TGN, they traffic through the Golgi apparatus to the endoplasmic reticulum (ER), where they are subsequently translocated through the Sec61p channel into the cytosol via ER quality control mechanisms (49, 56, 62). It is believed that this complex retrograde transport may allow for certain essential steps in toxin activation and transfer; Stx has been shown to be cleaved and activated by host proteases (16). Similarly, the ability to reach the ER may enable a chaperone-facilitated transfer to the cytosol, as previously reported for Ctx, ricin, and Stx (49, 52, 62, 63).

The existence of a retrograde transport pathway was first uncovered by electron microscopic studies tracking the intracellular transport of Stx (46). Since that time, a number of toxins, in addition to Ctx and ricin, have likewise been found to transit through the ER en route to the cytoplasm. Given the importance of these pathways to intoxication by diverse pathogenic agents, a number of investigations have been directed at identifying host molecules involved in toxin transport. Previous studies aimed at identifying essential components of the retrograde pathways of Stx, ricin, and Ctx have focused largely on the Rab family of small GTP-binding proteins. Members of the Rab family cycle between their GTP- and GDP-bound forms, which are related to their functions as regulators of vesicular traffic (53). Standard genetic approaches involving overexpression of dominant-negative mutants or small interfering RNA knockdowns of various Rabs have revealed the complexity of toxin trafficking pathways. Inhibition of Rab7 and Rab9, which are involved in lysosome targeting pathways, had no effect on ricin and Stx trafficking (21, 47). In contrast, inhibition of Rab6a′, involved in endosome-to-TGN transport, inhibited Stx transport from endosomes through the Golgi apparatus (17, 33, 59) but had no effect on ricin transport or intoxication (8). Similarly, overexpression of a dominant-negative Rab11, implicated in transport from recycling endosomes to the TGN, resulted in impaired Stx transport but had no effect on ricin (21, 60). Rab22, like Rab6a′, has been implicated in endosome-to-TGN transport, though inhibition of Rab22 function has had inconsistent effects on retrograde toxin transport (34). Though these pathways still remain poorly characterized, the sequential retrograde progression utilized by these toxins has translated into a unique system for probing host endocytic mechanisms.

In an effort to dissect and inhibit the stepwise trafficking of Stx, we have developed a quantitative and highly sensitive, high-throughput luciferase-based assay to screen a library of small-molecule compounds for their ability to block Stx-medi-
ated inhibition of protein synthesis (64). Because Stx transport involves a multistep progression through the cell, we predicted that inhibitory compounds could be identified at distinct stages along the retrograde trafficking pathway and could potentially be directed at specific molecular targets. From an initial screen of 14,400 small compounds, we identified several potential inhibitors. Among these, we characterized two compounds (compounds 75 and 134) that reversibly inhibit Shiga intoxication and act at distinct steps along the toxin trafficking pathway. Our results demonstrate the utility of a small-molecule approach to elucidating toxin transport pathways and will lead to the identification of novel therapeutic approaches targeting diseases caused by ER-routed toxins.

MATERIALS AND METHODS

Reagents and antibodies. Small chemical compounds were purchased from ChemDiv and reconstituted to 5-mg/ml stocks in dimethyl sulfoxide (DMSO). All reagents for purity testing were purchased from Sigma. Trans35S was purchased from MP Biomedicals, and Cycloheximide, DMSO, brefeldin A (BFA), nocodazole, anisomycin, and stau-rorosine (DMEM), Eagle's minimum essential medium, streptomycin, and penicillin were from Sigma. Recombinant Ctx subunit B (CtxB) labeled with Alexa Fluor 488, Alexa Fluor 594-labeled human transferrin (Tf), SlowFade Gold mounting reagent, and without or with 4,6-diamidino-2-phenylindole (DAPI), and Alexa Fluor-labeled goat or donkey secondary antibodies against immunoglobulin G were obtained from Molecular Probes. Rabbit anti-giantin was from Covance and sheep anti-human TGN46 from Serotec. Dulbecco's modified Eagle's medium (DME), Eagle's minimum essential medium, streptomycin, and penicillin were from BioWhittaker. Nonessential amino acids were purchased from Mediatech. Cyclomexidine, DMSO, brefeldin A (BFA), nacodazole, anisomycin, and stau-rorosine from Sigma. Tran35S was purchased from MP Biomedicals, and 35S-labeled O4 was obtained from American Radiolabeled Chemicals.

Cell culture. Vero cells were grown and maintained in DMEM supplemented with 10% fetal calf serum (Sigma), 100 μg/ml streptomycin, 100 U/ml penicillin, and 1% nonessential amino acids at 37°C under 5% CO2. Tran35S was added to wells of triplicates, and cells were shifted to 37°C for an additional 4 h. The medium was then removed from all wells and replaced with a medium containing Tran35S-label at 10 μCi/ml. Cells were incubated at 37°C for 45 min, washed with phosphate-buffered saline (PBS) (pH 7.4), and lysed (1 mg/ml bovine serum albumin, 0.2% deoxycholic acid, 0.1% SDS, 20 mM Tris [pH 7.4]) at 4°C for 12 h. Proteins from the lysed cells were precipitated with trichloro-acetic acid (TCA) (final concentration, 15%) and transferred to multiscreen HA plates (Millipore), and the filters were washed with ice-cold 20% TCA. Filters were then removed from the plate and placed in 2 ml BioSafe II scintillation fluid (RPI), and 35S incorporation was quantitated using a beta counter (Beckman). Independent experiments were performed at least three times for potent compounds, and data were analyzed using Prism software (version 4.0; 2003).

Toxin and Tf internalization. For toxin trafficking experiments, Vero cells were grown in chamber slides (2.5 × 104 cells/chamber), treated with a medium containing DMEM, compound, or known agents at the indicated concentrations for 1 h at 37°C, and then placed on ice for 15 min prior to the addition of toxin. The toxin was bound for 45 min at 4°C, followed by washing of unbound toxin with ice-cold PBS (pH 7.4). Fresh, prewarmed medium was added, and cells were shifted to 37°C for the indicated times to allow for toxin internalization. In Tf trafficking experiments, cells were pretreated with compounds in serum-free culture medium, and Tf and toxin were bound to cells at 4°C for 1 h, followed by a shift to 22°C for 1 h. For all immunofluorescence experiments, cells were fixed in 4% paraformaldehyde in cold PBS, permeabilized in a culture medium containing 1% Triton X-100, and washed with cold PBS containing 0.1% bovine serum albumin (wt/vol), all at room temperature. All primary antibodies and secondary antibodies (donkey anti-immunoglobulin G labeled with Alexa Fluor 488, 594, or 555) were diluted in blocking buffer. Cells were rinsed thoroughly in PBS prior to being mounted in SlowFade Gold reagent with or without DAPI (InVitrogen Corp.). Fluorescence imaging used epiillumination (Zeiss) or confocal (Olympus) microscopy.

Cell viability assays. The viability of cells treated with a compound was evaluated using the CellTiter-Glo luminescent cell viability assay (Promega), a luciferase-based assay. Vero cells (5 × 104/well) were added to 96-well plates and grown at 37°C under 5% CO2 overnight. The medium was then removed and replaced with a prewarmed medium either with DMEM alone or with a compound, in triplicate. Following incubation at 37°C for various times, an equal volume of CellTiter-Glo reagent (50 μl) was added according to the manufacturer’s instructions, and the light output was measured using the Lmax 1.1L luminometer. Independent experiments were performed three times.

Cloning and expression of StxB-Sulf2-His6. In order to add overlapping sulfation sites to the carboxyl terminus of StxB, the StxB gene from pNAS-13 (61) was amplified with primers Sulf1-5’-GGTGGCTACAAGTTGTGTAAT-3’ and Sulf2-5’-GGTGCTCAAGGAGTATTGTGTAAT-3’. The amplified product was ligated into expression plasmid pCRT7-TOPO (Invitrogen). The luciferase-based assay. Vero cells (5 × 104/well) were added to 96-well plates and grown at 37°C under 5% CO2 overnight. The medium was then removed and replaced with a prewarmed medium either with DMEM alone or with a compound, in triplicate. Following incubation at 37°C for various times, an equal volume of CellTiter-Glo reagent (50 μl) was added according to the manufacturer’s instructions, and the light output was measured using the Lmax 1.1L luminometer. Independent experiments were performed three times.

Large number of samples. Vero cells were cultured overnight at 37°C under 5% CO2 in 96-well plates at 2.5 × 104 cells/well, whereupon the medium was removed and replaced with either prewarmed medium (plus 0.5% [vol/vol] DMSO) or a medium containing a compound. Following a 1-h incubation at 37°C, toxin was added to wells in triplicate, and cells were shifted to 37°C for an additional 4 h. The medium was then removed from all wells and replaced with a medium containing Tran35S-label at 10 μCi/ml. Cells were incubated at 37°C for 45 min, washed with phosphate-buffered saline (PBS) (pH 7.4), and lysed (1 mg/ml bovine serum albumin, 0.2% deoxycholic acid, 0.1% SDS, 20 mM Tris [pH 7.4]) at 4°C for 12 h. Proteins from the lysed cells were precipitated with trichloroacetic acid (TCA) (final concentration, 15%) and transferred to multiscreen HA plates (Millipore), and the filters were washed with ice-cold 20% TCA. Filters were then removed from the plate and placed in 2 ml BioSafe II scintillation fluid (RPI), and 35S incorporation was quantitated using a beta counter (Beckman). Independent experiments were performed at least three times for potent compounds, and data were analyzed using Prism software (version 4.0; 2003).

Luciferase-based assay for measuring protein synthesis. The luciferase-based assay has been described previously (64). It was applied to a high-throughput screen (HTS) of small molecules consisting of known biological compounds as well as unknown compounds at the ICCB facility at Harvard University. The luciferase protein has been modified by Promega by the addition of a PEST sequence, resulting in its short intracellular half-life (44). The luciferase-PEST cDNA was cloned into an adenovirus expression plasmid (64), and high-titer viral stocks were generated (pAD-Luc). Vero cell monolayers were transduced with pAD-Luc (multiplicity of infection, 200), incubated for 24 h at 37°C under 5% CO2, and then seeded into 384-well black polystyrene plates (Corning) at 1 × 104 cells/well for an additional 24 h. Cells were then treated for 30 min at 37°C with a known biological compounds and unknown compounds at 5 ng/ml. Stx was added at 1 ng/ml, and cells were incubated for an additional 4 h at 37°C. To determine luciferase expression, the SuperLight luciferase reporter gene assay was used according to the manufacturer’s instructions (BioAssay Systems), and light output was detected using an LMax 1.1L luminometer ( Molecular Devices). To be considered protective, the compound must increase the luciferase signal at least twofold above the mean observed from cells treated with toxin alone.

Radioactive amino acid incorporation assay. Experiments measuring radioactive 35S incorporation were used to confirm positive hits from the luciferase-based HTS. This assay was adapted to a multowell format to enable testing on a large number of samples. Vero cells were cultured overnight at 37°C under 5% CO2 in 96-well plates at 2.5 × 104 cells/well, whereupon the medium was removed and replaced with either prewarmed medium (plus 0.5% [vol/vol] DMSO) or a medium containing a compound. Following a 1-h incubation at 37°C, toxin was added to wells in triplicate, and cells were shifted to 37°C for an additional 4 h. The medium was then removed from all wells and replaced with a medium containing Tran35S-label at 10 μCi/ml. Cells were incubated at 37°C for 45 min, washed with phosphate-buffered saline (PBS) (pH 7.4), and lysed (1 mg/ml bovine serum albumin, 0.2% deoxycholic acid, 0.1% SDS, 20 mM Tris [pH 7.4]) at 4°C for 12 h. Proteins from the lysed cells were precipitated with trichloroacetic acid (TCA) (final concentration, 15%) and transferred to multiscreen HA plates (Millipore), and the filters were washed with ice-cold 20% TCA. Filters were then removed from the plate and placed in 2 ml BioSafe II scintillation fluid (RPI), and 35S incorporation was quantitated using a beta counter (Beckman). Independent experiments were performed at least three times for potent compounds, and data were analyzed using Prism software (version 4.0; 2003).

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37°C for 1 h, fixed with 4% paraformaldehyde, and visualized by epifluorescence microscopy.

**Sulfation of Stx-B-Sulf-His.** Vero cells were seeded overnight at 37°C under 5% CO₂ (1 × 10⁶ cells/well). The next day, the medium was replaced with serum-free DMEM lacking sulfate (Washington University Tissue Culture Support Center), and cells were incubated for an additional 3.5 h at 37°C. The medium was replaced with sulfate-free DMEM containing DMSO or compound (for 30 min at 37°C); then it was replaced with prewarmed sulfate-free medium containing 1 mM/μl 35S-labeled O4 for 3 h at 37°C. Wells were washed with cold PBS (pH 7.4) and lysed with PBS containing 1% Triton X-100. The protein concentrations of postnuclear supernatants were determined by the bicinchoninic acid protein assay (Fierce), and 250 μg of lysates was added to 40 μl of nickel-nitritotriacetic acid Superflow beads and rotated at 4°C overnight. The next day, beads were spun down at 5,000 rpm for 5 min, and the unbound fraction was collected. Total 3S incorporated counts were determined by measuring radioactive counts from TCA-precipitated proteins of unbound lysates. Beads were washed once with PBS containing 1% Triton X-100 and twice with PBS. Beads were resuspended in imidazole (1.5 M in PBS), and eluates were denatured with 1× SDS gel-loading buffer (50 mM Tris-HCl, 100 mM β-mercaptoethanol, 2% SDS, 0.1% bromphenol blue, 10% [vol/vol] glycerol) and boiled. Eluates were resolved on a 10 to 20% Tris-HCl denaturing gel, fixed, and developed overnight in a phosphorimager cassette.

**Expression and trafficking of VSVG-GFP.** Vero cells were transiently transfected with vesicular stomatitis virus G protein (VSVG)-green fluorescent protein (GFP) at 0.045, as previously described for other cell lines (19). Briefly, 10⁶ Vero cells were transfected with 20 μg of pCDM8.1 expressing VSVG-GFP and a Lipofectamine 2000 (Invitrogen) mixture in antibiotic-free medium (DMEM with 10% fetal calf serum) followed by overnight incubation at 37°C under 5% CO₂. Cells were collected and placed in chamber slides (Lab-Tek) for an additional 8 to 10 h at 37°C before their transfer to 42°C for 12 to 16 h. Cells were then treated with compounds or BFA (25 μg/ml) in prewarmed, antibiotic-free medium for 1 h at 42°C before being transferred to 32°C. Thirty minutes prior to the shift to 32°C, all chambers were treated with cycloheximide (100 μg/ml) to prevent de novo protein synthesis. Cells were fixed following various incubation times at 32°C. Fixation, permeabilization, staining, and imaging were performed as described for toxin and TF internalization experiments.

**GFP secretion time course.** A plasmid encoding enhanced GFP (EGFP) fused to the amino terminus with the secretion signal of human neutrophile Y (NPY) was obtained from Richard Mains (12). The insert was released by BglII and NotI digestion and ligated into plasmid pENTR-4 (Invitrogen) that had been digested with BamHI and NotI. This plasmid was used as the source for insertion of the NPY-EGFP DNA into adenovirus expression plasmid pAD-DEST (Invitrogen) by the clonase reaction (Invitrogen). The resulting plasmid, pAD-NPY-GFP, was linearized with PciI and transfected into 293A cells to prepare a low-titer viral stock. The virus was amplified by further passage in 293A cells until a high-titer stock was prepared. Vero cells transduced with pAD-NPY-GFP were washed, trypsinized, and seeded into 6-well chamber slides. The next day, the medium was replaced with a culture medium containing DMSO or the indicated compound for 30 min at 37°C. Cycloheximide was then added at 100 μg/ml, and cells were washed and fixed with 4% paraformaldehyde in PBS (pH 7.4) at various times following cycloheximide treatment. Permeabilization, staining, and imaging were performed as described for toxin and TF internalization experiments.

**Assessment of NPY-GFP secretion.** Approximately 5 × 10⁶ Vero cells were infected overnight at 37°C under 5% CO₂ with pAD-NPY-GFP. Cells were then washed, trypsinized, and seeded into each well of a 6-well plate (1 × 10⁶ cells/well). The next day, the medium was removed, and cells were washed twice with serum-free medium. The medium was subsequently replaced with serum-free DMEM containing DMSO or a compound for 30 min at 37°C. Cells were all treated with cycloheximide (100 μg/ml) to inhibit de novo protein synthesis and to synchronize NPY-GFP trafficking. At various time points following cycloheximide treatment, supernatants were collected, and GFP secretion was assessed by an enzyme-linked immunosorbent assay (ELISA) as described in the manufacturer’s instructions (Pierce). ELISA plates were analyzed by the GenS software program (BioTek) using the Synergy 2 spectrophotometer (BioTek). The mean absorbance for control wells containing DMEM alone was subtracted from the absorbance for each sample well before analysis.

**Statistics.** All statistical analyses were performed by GraphPad Prism 5. For Fig. 1, toxin concentrations were log transformed prior to curve fitting and statistical analyses. Toxicity-response curves were generated by nonlinear regression (least-squares fit) to correspond to the observed data, and the concentration of toxin needed to reduce protein synthesis by 50% (50% inhibitory concentration [IC₅₀]) was calculated by using the fitted curves. For the toxicity-response curves, the toxin concentration varied, while the concentration of DMSO (0.5% [vol/vol] in control cells), compound 75 (25 μM), or compound 134 (50 μM) was kept constant. Toxic IC₅₀s were compared using the extra sum-of-squares F test applied to the best-fit curves for the data. Differences between toxin IC₅₀s were considered statistically significant at a P value of ≤0.05 and highly statistically significant at a P value of ≤0.01.

**RESULTS**

**HTS for compounds that inhibit Stx activity in host cells.** Several toxins damage host cells by inhibiting protein synthesis. DT and *Pseudomonas* exotoxin inhibit protein synthesis through the ADP-riboseylation of elongation factor 2 (7, 41), whereas Stx and the plant toxin ricin inhibit ribosome function by cleaving an adenosine residue from the 60S ribosome (3, 13, 39, 40, 45). Still other toxins, such as Ctx and anthrax edema toxin, induce increases in second-messenger levels, resulting in cytotoxicity (6, 26, 54). In order to quantitate the effects of various protein synthesis-inhibiting toxins on host cells, we had previously established a luciferase-based assay that could readily determine the susceptibilities of various cell lines to Stx, ricin, DT, and *Pseudomonas* exotoxin (64). In cells constitutively expressing an mRNA encoding destabilized firefly luciferase, luciferase enzyme activity served as a surrogate measure of protein synthesis.

This assay was adapted to an HTS and applied to a screen of small-molecule compounds that inhibit toxin susceptibility. The ICCB facility at Harvard University contains a number of commercial libraries consisting of synthetic and natural products. An initial screen of biological compounds with known effects yielded positive hits such as BFA and di-h-threo-1-phe-nyl-2-decanoylamino-3-morpholinol-1-propanol (PDMP) (data not shown), two compounds previously shown to inhibit Stx susceptibility through distinct mechanisms (25, 50) and serving as positive controls for the detection of Stx-inhibitory compounds. In addition, the assay detected known inhibitors of the proteasome, such as MG-132 (3). This was an expected result, since the discriminatory power of the assay is dependent on the rapid degradation of luciferase following translation (65).

We next screened the ChemDiv 3 library at the ICCB facility, consisting of 14,400 compounds of unknown function. The compounds included in this library were selected for their structural diversity, chemical stability, and “drug-like properties.” Since these compounds were commercially available and their functions currently undefined, we reasoned that novel inhibitors could be identified. Among these were selected the top 1% of compounds yielding the highest luciferase signal in the presence of toxin, all of which resulted in a signal at least twice the baseline. Because the initial screen lacked a counter-screen to exclude compounds affecting luciferase turnover, each compound was subsequently tested for its effect on the luciferase signal following cycloheximide treatment. Cyclohexi-mide-mediated inhibition of protein synthesis is independent of intracellular transport, and its mechanism of ribosomal inactivation is distinct from that of Stx (55). Therefore, compounds that provide protection against cycloheximide-mediated inhibition of the luciferase signal must be acting in a toxin-independent manner (e.g., by inhibiting luciferase degradation), and such compounds were excluded from further analysis. After exclusion of compounds that affected cycloheximide-induced suppression of the luciferase signal, eight com-
data point (mean \pm standard deviation) represents triplicate data at the indicated toxin concentration from one representative experiment.

FIG. 1. Protective effects of inhibitory compounds against Stx, ricin, and DT. (A) Compound 134 structure and demonstration that this compound (50 \mu M) protects against Stx- and ricin- but not DT-mediated decreases in protein synthesis. Toxin IC50s for Stx and ricin in compound-treated cells was found to be statistically different (\( P < 0.05 \)) from those in control cells. The DT IC50 in compound-treated cells was not statistically different from that in control cells (\( P > 0.05 \)). (B) Compound 75 structure and demonstration that this compound (25 \mu M) protects against Stx-, ricin-, and DT-mediated decreases in protein synthesis. Toxin IC50s for all three toxins in compound-treated cells were found to be statistically different (\( * P < 0.01 \)) from those in control cells. The DT IC50 in compound-treated cells was found to be highly statistically different (\( ** P < 0.001 \)) from those in control cells. The DT IC50 in compound-treated cells was found to be statistically different (\( *** P < 0.0001 \)) from those in control cells.

Secondary analysis to determine the potency and efficacy of inhibitory compounds. The optimal protective concentration for each identified hit was determined using a radioactive assay for protein synthesis (11) that was modified for medium-throughput analysis in a multiwell format (see Materials and Methods). Since some of these compounds could exhibit non-specific effects at increased concentrations, the lowest concentration providing significant protection against Stx compared to the effect of the toxin on untreated Vero cells was considered to be optimal. Compounds classified as inhibitors showed half-maximal activity between 10 and 50 \mu M (data not shown). Compounds were used above their half-maximal but below their maximal concentrations for all subsequent assays (at 25 \mu M for compound 75 and 50 \mu M for compound 134).

The abilities of these compounds to protect against increasing Stx concentrations were expressed as the toxin IC50s (see Materials and Methods). Using these criteria, compounds 75 and 134, at their respective optimal concentrations, exhibited the greatest protective effects among the hits identified from the initial screen. Both compounds showed statistically significant increases in the Stx IC50 compared to that for cells containing no compound (Fig. 1). At higher concentrations (50 \mu M and 100 \mu M, respectively), compounds 75 and 134 exhibited up to 1,000-fold increases in the Stx IC50 (data not shown).

Neither of these compounds affected luciferase degradation in the presence of cycloheximide (data not shown).

An initial characterization of compounds showing highly protective effects against Stx led us to consider whether these compounds could protect against other toxins that inhibit protein synthesis. Compounds 75 and 134 showed similar statistically significant increases in the ricin IC50 (Fig. 1) (\( P < 0.01 \)). Protection against both Stx and ricin was also greater than the previously observed protective effects of an overexpressed dominant-negative mutant of Rab2, a small GTP-binding protein found to be essential to Stx transport through the Golgi apparatus (8).

Interestingly, compound 134 failed to show a statistically significant effect against DT-mediated protein synthesis inhibition (Fig. 1A), while compound 75 demonstrated greater protection (Fig. 1B). Vero cells that were not treated with a compound demonstrated a DT susceptibility profile similar to those for Stx and ricin. Stx and ricin, following endocytosis, are known to traffic from an endosomal compartment to the ER via the Golgi apparatus (30). DT, however, directly accesses the cytosol from early endosomes; the low endosomal pH is believed to allow for a conformational change in the holotoxin and to promote shuttling of the A moiety across the endosomal membrane (41). The lack of protection against DT suggests that compound 134 affects toxin transport at a point after the early-endosome stage but has no effect on trafficking from the plasma membrane to the early endosomal compartment. In contrast, the protective effect of compound 75 against all three toxins suggests that this compound affects transport mecha-
nisms common to all three pathways, presumably at an earlier stage in endocytosis than the site of action of compound 134.

**Effects of compounds 75 and 134 on toxin transport.** In order to determine the site at which inhibitory compounds were affecting toxin activity, the endocytosis and transport of fluorescent StxB and CtxB were examined. The retrograde transport of protein toxins is believed to occur exclusively from early and/or recycling endosomes (32). In order to determine whether endocytosis and trafficking of StxB to early and recycling endosomes were affected by compound 134, StxB transport was compared to that of fluorescently labeled Tf, which is known to accumulate in recycling endosomes at 22°C due to a block in recycling-endosome-to-TGN transport at this temperature (33). In agreement with previous reports, StxB colocalized with Tf-positive compartments in control cells at 22°C (Fig. 2A). StxB trafficking to Tf-positive compartments in compound 134-treated cells was not significantly different from that in control cells, showing a similar level of colocalization with the perinuclear recycling-endosome compartment (Fig. 2A). Similar results were seen with CtxB, which also accumulated in a Tf-positive perinuclear compartment. (Fig. 2B). In addition, compound 134 did not affect the binding of StxB to the cell surface (see Fig. S2 in the supplemental material), suggesting that this compound was not occupying toxin receptor binding sites or significantly decreasing receptor expression. Taken together with the relative lack of protection by compound 134 against DT-mediated protein synthesis inhibition (Fig. 1A), these results collectively suggest that compound 134 maintains Stx and Ctx transport to recycling endosomes.

Compound 75, like compound 134, did not inhibit StxB binding to its receptor or decrease receptor expression (see Fig. S2 in the supplemental material). However, treatment with compound 75 appeared to inhibit the transport of both StxB/CtxB and Tf to perinuclear recycling endosomes. Most of the toxin and most of the Tf were located in peripheral vesicular structures (Fig. 2A and B), likely early endosomes. These results, in addition to those demonstrating the effects of compound 75 on susceptibility to DT, collectively suggest that this compound inhibits Stx and Ctx transport to recycling endosomes.

Immunohistochemical analysis of compound-treated cells revealed morphological changes to the Golgi apparatus (see Fig. S3 in the supplemental material). The dispersal of the Golgi apparatus is a morphological effect that has been observed under several conditions that impair retrograde and intra-Golgi transport (10). To exclude the possibility that the effects of these compounds were not specific to intracellular toxin transport, we sought to rule out processes that are known to have effects on Golgi morphology. In particular, during the process of apoptosis, the Golgi apparatus becomes fragmented (9, 31, 43). In order to rule out the possibility that compounds 75 and 134 were inducing Golgi apparatus fragmentation through apoptosis, ATP levels in compound-treated cells were compared to those in untreated cells or cells treated with staurosporine, a known apoptosis-inducing agent (23). Treatment with compound 75 or 134 failed to deplete ATP levels over 24 h, in contrast to the effect observed following staurosporine treatment, suggesting that the compounds were not cytotoxic or apoptosis inducing at the specified concentrations over the time course studied (Fig. 3A). In addition, compound-treated cells consistently showed a more punctate and swollen Golgi apparatus compared to the more tubulated Golgi apparatus in staurosporine-treated cells (Fig. 3B). The Golgi morphology of compound-treated cells also differed from the characteristic Golgi architecture of cells treated with anisomycin, a peptidyl transferase inhibitor shown to induce apoptosis in

![FIG. 2. Compound 75 impedes StxB/CtxB and Tf ligand trafficking to recycling endosomes.](image)
small-molecule inhibitor of toxin transport. Small-molecule compounds may act by reversible or...
irreversible mechanisms. In general, irreversible inhibitors act by covalent interaction with or modification of target molecules, such as through oxidation or acylation. Reversible compounds often act as competitive inhibitors of enzymatic activity or of protein-target interactions. Morphological effects on the Golgi apparatus produced by compound 75 or compound 134 treatment were found to be reversible. Following washout of the compounds, the Golgi apparatus reassembled in compound-treated cells at a time point at which BFA-treated cells maintained an altered Golgi morphology (Fig. 5A). More importantly, Vero cells lost protection against Stx following washout of either compound (Fig. 5B).

**Effects of compounds on anterograde transport.** The toxin-trafficking pathway, proceeding from an early endosome to the ER via the Golgi apparatus, is almost the mirror inverse of the biosynthetic and secretory pathways. The degree of overlap between the retrograde and anterograde pathways, however, remains largely unresolved. Since no host molecules have been shown to traffic from endosomes all the way to the ER, the ability to track bacterial toxins and identify compounds that potentially block their transport allows us to specifically probe the retrograde pathway in its entirety.

To study the effects of these compounds on general anterograde transport, the sequential trafficking of a temperature-sensitive VSVG-GFP (tsf45) fusion protein (19) in transiently transfected Vero cells was examined. Cells kept at 42°C showed that VSVG-GFP was confined to the ER, and a shift to 32°C allowed anterograde progression to the Golgi apparatus by 1 h (Fig. 6). Compounds 75 and 134 did not inhibit VSVG-GFP transport from the ER to the Golgi apparatus, as evidenced by colocalization of VSVG-GFP with TGN46 as early as 1 h following the shift to 32°C (Fig. 6). BFA has been shown to impede ER-to-Golgi apparatus transport (42), and cells treated with BFA showed a dispersed Golgi apparatus, with VSVG-GFP restricted to the ER, over the time course studied. We conclude that both compounds 75 and 134 do not inhibit ER-to-Golgi apparatus transport of VSVG-GFP, suggesting that generalized anterograde transport mechanisms from the ER to the Golgi apparatus remain unaffected.

To equally monitor compound effects on the secretory pathway beyond the Golgi apparatus, secretion of a GFP construct bearing an amino-terminal NPY secretion signal (NPY-GFP) was assessed qualitatively and quantitatively. During the process of secretion, a portion of NPY-GFP is maintained in a compartment adjacent to and overlapping with the TGN (data not shown). In contrast to BFA and compound 75, compound 134 had a minimal effect on NPY-GFP secretion as judged by the immunofluorescence time course (see Fig. S4 in the supplemental material). Compound 75 partially impeded GFP secretion from the TGN-associated compartment, as evidenced by the persistence of GFP fluorescence as long as 1 h following cycloheximide treatment, at which time no GFP-positive cells were seen among control cells (see Fig. S4 in the supplemental material). An ELISA was developed to quantify the amounts of NPY-GFP secreted into the medium at various time points following cycloheximide treatment (to inhibit new protein synthesis). Mean absorbance values for control wells containing DMEM alone were subtracted from absorbance values for sample wells. Using this assay, we measured the effect of compound 134 (50 μM) on GFP secretion. The level of GFP secretion by compound 134-treated cells was 86% of that by control (untreated) cells at 1 h and 89% of that by control cells at 2 h. (data not shown). Compound 75 (25 μM) decreased post-Golgi secretion of NPY-GFP to 58% and 71% of that by control cells at 1 and 2 h, respectively (data not shown).

**DISCUSSION**

Using a luciferase-based assay that was adapted to an HTS for determining cell susceptibility to toxin-mediated inhibition of protein synthesis, we have screened a chemical library of small compounds and characterized two compounds that showed marked potency and selectivity against intracellular toxin transport. Both compounds demonstrated strong protective effects against Stx and ricin, and compound 75 was shown to protect equally against DT. We thus report two novel compounds demonstrating efficacy against multiple bacterial toxins, with toxin IC₅₀'s nearly an order of magnitude greater than those reported previously using knockdown approaches for toxin inhibition (1, 8). More importantly, our results demonstrated that compounds 75 and 134 conferred toxin-protective effects by disrupting transport at distinct steps along the toxin-trafficking pathway.

Screens at the ICCB facility have recently identified small-molecule inhibitors of *Toxoplasma gondii* invasion and *Vibrio cholerae* virulence. From a library of 12,160 compounds, 24 inhibitors of *Toxoplasma gondii* invasion were identified (5). These included compounds that inhibited host uptake of the organism as well as inhibitors of *Toxoplasma* gliding motility and microneme-based secretion. Effective doses of these compounds ranged from 3 to 100 μM. Recently, a similar screen identified a small-molecule inhibitor of *Vibrio cholerae* virulence (20). A total of 50,000 compounds were screened, and 109 compounds that inhibited virulence factor expression were identified. Of these, a compound named virastatin inhibited the transcriptional regulator ToxT, thereby suppressing the
expression of Ctx and the toxin-coregulated pilus. The MIC of virastatin against Ctx expression ranged from 3 to 40 μM, depending on the bacterial strain. Thus, recent screens underscore the utility of small-molecule assays in the identification of compounds that inhibit microbial virulence or block intracellular trafficking pathways.

Morphological analysis of compound-treated cells revealed dispersed Golgi apparatus-derived vesicles, but their pronounced toxin-protective effects were distinct from those of apoptosis-inducing or cytotoxic agents. The disrupted Golgi membranes showed competent anterograde trafficking of VSVG-GFP and secretion of NPY-GFP in compound 134-treated cells, implying that this compound was preferentially targeting components of the retrograde pathway. Similar Golgi apparatus fragmentation has been observed in HeLa cells depleted of Cog3p, and the disrupted Golgi membranes were equally capable of anterograde trafficking of VSVG (65). Though it also exhibited a strong protective effect against toxin-mediated protein synthesis inhibition, compound 75 appeared to have a more pronounced effect on the post-Golgi trafficking of NPY-GFP. Nonetheless, the identifica-
tion of two compounds preferentially targeting the retrograde pathway could represent a useful tool in elucidating similarities and differences between anterograde and retrograde trafficking.

The retrograde transport of protein toxins is believed to occur exclusively from early and/or recycling endosomes (33), and it would be tempting to assume that toxins such as Stx, Ctx, and ricin subvert retrograde sorting pathways utilized by certain host proteins, such as acid hydrolase receptors (4), for endosome-to-TGN transport. Studies increasingly show that protein toxin transport to the TGN is not uniform (48), underscoring the complexity of retrograde pathways leading to a similar destination. A previous study examining toxin transport found that disruption of the Golgi apparatus by the expression of a temperature-sensitive mutant of ε-toxin lectins on eukaryotic ribosomes. Nucleic Acids Symp. Ser. 2003. Evidence that the toxoplasma exotoxin contains a specific sequence at the carboxyl terminus that is required for cytotoxicity. Proc. Natl. Acad. Sci. USA 101:2577–2580.


