Discovery and Analysis of 4H-Pyridopyrimidines, a Class of Selective Bacterial Protein Synthesis Inhibitors

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Bacterial protein synthesis is the target for numerous natural and synthetic antibacterial agents. We have developed a poly(U) mRNA-directed aminoacylation/translation protein synthesis system composed of phenyl-tRNA synthetases, ribosomes, and ribosomal factors from Escherichia coli. This system, utilizing purified components, has been used for high-throughput screening of a small-molecule chemical library. We have identified a series of compounds that inhibit protein synthesis with 50% inhibitory concentrations (IC50s) ranging from 3 to 14 μM. This series of compounds all contained the same central scaffold composed of tetrahydropyrido[4,3-d]pyrimidin-4-ol (e.g., 4H-pyridopyrimidine). All analogs contained an ortho pyridine ring attached to the central scaffold in the 2 position and either a five- or a six-member ring tethered to the 6-methylene nitrogen atom of the central scaffold. These compounds inhibited the growth of E. coli, Staphylococcus aureus, Streptococcus pneumoniae, Haemophilus influenzae, and Moraxella catarrhalis, with MICs ranging from 0.25 to 32 μg/ml. Macromolecular synthesis (MMS) assays with E. coli and S. aureus confirmed that antibacterial activity resulted from specific inhibition of protein synthesis. Assays were performed for the steps performed by each component of the system in order to ascertain the target of the compounds, and the ribosome was found to be the site of inhibition.

Bacterial infections continue to represent a major worldwide health problem. Infections range from the relatively innocuous, such as skin rashes and common ear infections in infants, to serious and potentially lethal infections in immune-compromised patients. Resistance to antibacterial agents has increased in many pathogenic bacteria and can occur through a variety of mechanisms, such as target mutation, induction of efflux pumps, or induction of metabolic pathways leading to the degradation of the compound. Resistance developed in one cell can be transferred to other bacteria by horizontal gene transfer. The need for new antibiotics to address the increase in resistance has become critical.

Antibacterial agents interfere with cellular processes that are essential for the survival of the cell (for a complete list, see reference 4). For both naturally occurring and synthetic antibiotics, protein synthesis is a major target of antibiotic action. Bacterial protein synthesis inhibitors include the macrolides (e.g., erythromycin, clarithromycin, and azithromycin), clindamycin, chloramphenicol, the aminoglycosides (e.g., streptomycin, gentamicin, and amikacin), and the tetracyclines (2, 18, 49). The newest class of antibacterials, the synthetic oxazolidinones (exemplified by linezolid, the only novel and approved ribosomal inhibitor), also inhibit protein synthesis (21, 45). Protein synthesis is the cellular process most frequently targeted by naturally occurring antibacterials, providing compelling evolutionary evidence for the susceptibility of this process to antibiotic intervention (21).

There has been much well-deserved ado recently concerning access to the crystal structures of ribosomes, either alone or bound to a variety of antibiotics (2, 17, 35, 44). This work has led to great progress in refining the effectiveness of these classes of inhibitors via structure-based drug design (19). However, extant resistance mechanisms may also be circumvented by identifying new structural classes that bind in substantially different ways or at different sites on the ribosome. In addition, certain molecular inhibitors bind to and inhibit their targets in an induced-fit mode (12), and this has been seen with some ribosomal inhibitors (10). Since this type of interaction may not be immediately recognized in a structure-based design process, the discovery of inhibitors of function remains a useful method for novel drug discovery.

The ribosome is a well-established target for drug discovery, but other components that are essential for protein synthesis also offer attractive targets. Elongation factor Tu (EF-Tu) delivers the charged tRNA to the A-site of the ribosome in a ternary complex with GTP and an aminoacylated tRNA, hydrolyzing the GTP to GDP in the process (14, 39). Elongation factor Ts (EF-Ts) then interacts with EF-Tu to regenerate EF-Tu to an active form, facilitating the replacement of bound GDP with GTP (50). Elongation factor G (EF-G) plays a central role in the elongation phase of protein synthesis by catalyzing GTP-dependent translocation (1, 13, 40). EF-G is also one of the proteins involved in the termination of protein synthesis in a GTP-dependent fashion (47). Amino-acyl tRNA synthetases (aaRS) catalyze
the attachment of amino acids to their cognate tRNAs. They are essential components in protein synthesis and individually provide attractive targets for the discovery of antibiotics (42).

Recently, attempts have been made to screen chemical-compound libraries by using cell extracts containing native transcription and translation systems from *Escherichia coli* (37), *Streptococcus pneumoniae* (9, 37), and *Staphylococcus aureus* (28). This approach has had only limited success. The use of cell extracts for screening can be problematic due to the presence of nucleases, degraded nucleic acids, soluble but denatured proteins, and turbidity (22). In addition, different preparations of S30 fractions can differ in activity and are therefore undependable (23). To avoid these problems, we have developed a polyclonal-directed aminoacylation/translation (AT) protein synthesis system composed of phenyl-tRNA synthetases, ribosomes, and ribosomal factors from *E. coli*. Using this system as a platform for screening, we have discovered a compound series capable of inhibiting protein synthesis in vitro and in whole-cell assays. The development of the screening system and the characterization of the resulting inhibitors is described.

**MATERIALS AND METHODS**

**Gel electrophoresis and protein analysis.** Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) was performed using either 12% or 4 to 12% polyacrylamide precast gels (Novex NuPAGE; Invitrogen) with morpholinepropanesulfonic acid (MOPS) running buffer (Invitrogen). Photographs and densitometry were performed using a Kodak Image Station, model 440CF. Protein concentrations were determined by the method of Bradford (3) using bovine serum albumin as the standard.

**Purification of ribosomes and proteins.** Early-phase ribosomes from *E. coli* strain MRE600 were prepared in the Hill laboratory at the University of Montana (Missoula) as previously described (46). Native *E. coli* EF-Tu was purified from cells grown to an optical density of 2.0. The cells were first lysed with a homogenizer (Niro) and then clarified by centrifugation (22,000 g) at 4°C and the protein was precipitated using ammonium sulfate. The protein precipitating between 35 and 75% saturated ammonium sulfate was collected by centrifugation (23,000 × g, 60 min, 4°C) and further purified using DEAE ion exchange (GE Healthcare) and Superdex-200 (GE Healthcare) size exclusion chromatography on an AKTA liquid chromatography system (GE Healthcare). The resulting protein was more than 98% homogeneous.

The gene encoding *E. coli* EF-G was PCR amplified from *E. coli* genomic DNA using forward primer 5′-CACCAGTCATCAGCAGCATCGCAGGCGTCGT -3′ and reverse primer 5′-CTATATTACATACGGGCTTCAA, which was designed to add six histidine amino acid residues to the N terminus. The PCR product was inserted into a pET101D/TOPO vector (Invitrogen), transformed into Rosetta (DE3) (Novagen) cells, and expressed as an N-terminally His tagged protein. EF-G was purified to more than 98% homogeneity using nickel-nitrilotriacetic acid (NTA) affinity chromatography (Qiagen).

The genes encoding the *E. coli* PhrEα and β subunits were PCR amplified from genomic DNA as a new operon using 5′-CACCAGTCATCAGCAGCATCGCAGGCGTCGT -3′ and 5′-ACTATATCATCAGCAGCATCGCAGGCGTCGT -3′ as a forward primer and 5′-ACTATATCATCAGCAGCATCGCAGGCGTCGT -3′ as a reverse primer, and the PCR product was inserted into pET101D/TOPO (Invitrogen). The resulting plasmid was transformed into BL21 Star (DE3) cells (Novagen), and the two subunits were expressed in their native forms. The cells were lysed; the lysate was clarified by centrifugation; and the protein was precipitated using ammonium sulfate (as described above). The proteins precipitating between 40 and 55% saturated ammonium sulfate were further purified using DEAE ion exchange and Superdex-200 size exclusion chromatography. PhrEα was purified to more than 98% homogeneity.

Bacteria overexpressing an N-terminally His tagged form of *E. coli* EF-Ts were a gift from the Spremulli laboratory (University of North Carolina, Chapel Hill). EF-Ts was purified to more than 98% homogeneity using Ni-NTA affinity chromatography (Qiagen).

**AT assays.** A scintillation proximity assay (SPA) was developed for the AT assay. The complete assay mixture contained 50 mM Tris-HCl (pH 7.5), 40 mM KCl, 10 mM MgCl₂, 0.1 mM spermine, 1.5 mM ATP, 0.5 mM GTP, 25 μM [³H]phenylalanine (100 cpm/pmol), and 0.25 mg/ml poly(U). To maintain constant levels of ATP and GTP, the assay mixture contained a nucleotide regeneration system composed of 4.75 mM phosphoenolpyruvate (PEP) and 0.026 U/μl pyruvate kinase (PK). The concentrations of ribosomes and proteins in the assay were as follows: ribosome, 0.11 μM; PhrEα, 0.025 μM; EF-Ts, 0.9 μM; EF-Ts, 0.03 μM; EF-G, 0.16 μM. These concentrations were arrived at through sequential rounds of optimization: each concentration is just below the saturation point of the titration.

The screening reactions were carried out in 96-well microtiter plates (Costar). Test compounds were equilibrated by the addition of 39 μl of the protein-substrate mixture (without tRNA) to 1 μl of the chemical compound (3.2 mM) dissolved in 100% dimethyl sulfoxide (DMSO). This mixture was allowed to incubate at ambient temperature for 15 min, and then reactions were initiated by the addition of 10 μl of *E. coli* RNA (150 μM), followed by a 2-h incubation at room temperature (comparable to 1 h at 37°C). Reactions were stopped by the addition of 5 μl of 0.5 M EDTA. Two hundred micrograms of SPA beads (RNA binding beads [Ysi]; Perkin-Elmer) in 150 μl of 300 mM citrate buffer (pH 6.2) was added. The plates were analyzed using a Packard Topcount NXT scintillation counter. Assays to determine 50% inhibitory concentrations (IC₅₀) were carried out as described above with the test compounds serially diluted from 200 μM to 0.39 μM. The concentration ranges of antibiotics in control plates were as follows: spiramycin, 0.07 μM to 18.0 μM; tylosin, 0.05 μM to 13.0 μM.

**PhrEα assay.** SPA5s to determine PhrEα inhibition by chemical compounds were carried out as described previously (6), with the exception that the enzyme mixture was preincubated with 0.4 to 200 μM compound for 15 min prior to the addition of tRNA. The reactions were stopped by the addition of 5 μl of 0.5 M EDTA. Four hundred micrograms of SPA beads (polyethyleneimine [PEI]-polivynil toluene [PVT] beads; Perkin-Elmer) in 150 μl of 300 mM citrate buffer (pH 2.0) was added, and the plates were analyzed as described above.

**EF-Tu GDP exchange assay.** Nitrocellulose binding assays were used to determine inhibition of GDP exchange by EF-Tu as previously described (5, 38), with the exception that 0.75 μM EF-Tu was preincubated with 0.4 to 200 μM compound for 15 min prior to the addition of [³H]GDP. The reactions were stopped by the addition of 5 μl of 0.5 M EDTA. Four hundred micrograms of SPA beads (polyethyleneimine [PEI]-polivynil toluene [PVT] beads; Perkin-Elmer) in 150 μl of 300 mM citrate buffer (pH 2.0) was added, and the plates were analyzed as described above.

**EF-Tu ternary-complex formation assay.** In a ternary complex, the acceptor stem and attached amino acid are protected by EF-Tu from hydrolysis by RNase A (24). Inhibition of EF-Tu in ternary-complex formation and in the protection of tRNA from hydrolysis was analyzed using filter binding assays as previously described (5). Enzyme mixtures contained 6 μM EF-Tu and 1.5 mM [³H]PhetRNA, and were preincubated with 0.8 to 400 μM compound for 15 min at 37°C prior to digestion with RNase A.

**Ternary-complex-ribosome binding assay.** Aminoacylated tRNAs are delivered to the A-site on the ribosome in a ternary complex composed of EF-Tu, GTP, and Phe-tRNA⁷. To determine ternary-complex binding to the ribosome, a mixture similar to the A/T assay mixture was used except that phenylalanine, PhrEα, and ATP were removed, and deacylated tRNA was replaced by [³H]Phe-tRNA⁷. GTP was also replaced with the nonhydrolyzable analog guanosine 5′-[β,γ-imido]triphosphosphate (GDPNP). All compounds except for the ribosomes were preincubated with 0.8 to 400 μM compound at 37°C for 15 min. Ribosomes were then added (0.5 μM), and incubation was continued for an additional 15 min. Assay products were analyzed using glass microfiber filter binding (Whatman) as previously described (7).

**EF-G GTPase assay.** Assay mixtures for ribosome-dependent GTP hydrolysis by EF-G contained 50 mM Tris-HCl (pH 7.5), 10 mM MgCl₂, 70 mM NH₄Cl, 1 mM dithiothreitol (DTT), 1.8 mM GDP, 0.04 μM ribosomes, and 2.2 μM EF-G. Mixtures were assembled on ice, and 48 μl of the mixture was added to 2 μl of the compound, transferred to 37°C, and incubated for 30 min. The final concentration of the compound ranged from 0.8 to 400 μM. The reaction was stopped by spotting 6.0 μl onto PEI-cellulose plates (Selecto Scientific). GTP and PP, were separated by thin-layer chromatography (TLC) using 4 M urea-1× TCEP (pH 3.5) as a mobile phase (26). GTP, PP, and P were quantified by phosphorimaging with a Storm 840 phosphorimag.
components were assembled on ice, mixed with the compound, and incubated at ambient temperature for 2 h. Reactions were stopped by the addition of 2 ml of 5% trichloroacetic acid (TCA), and reaction products were heated to 90°C for 15 min and filtered through glass fiber filters (Whatman). The concentrations of the compound ranged from 0.3 to 300 μM, and the concentration of the control, cycloheximide, ranged from 0.3 to 300 μM.

**Microbiological assays.** Broth microdilution MIC testing was performed in 96-well microtiter plates according to National Committee for Clinical Laboratory Standards (NCCLS; now CLSI) document M7–A6 (30). MICs were determined for *E. coli* toLC mutants, *E. coli* ATCC 25922, polymyxin B nonapeptide (PMBN)-treated *E. coli*, *S. aureus*, and *Streptococcus pneumoniae* as previously described (31). Secondary tests were carried out against *Hemophilus influenzae*, *Enterococcus faecalis*, and *Moraxella catarrhalis*.

Macromolecular synthesis (MMS) assays were performed in cultures of *E. coli* toLC mutants and in cultures of wild-type *S. aureus* as described previously (31). Briefly, *E. coli* ΔtolC CGSC5633 was obtained from the *E. coli* Genetic Stock Center (Yale University). This strain is an efflux mutant and was used to evaluate the global mode of action of the hit compounds. Assays were performed using the radiolabeled precursors [3H]thymidine, [5-3H]uridine, and L-[4,5-3H]the radiolabeled precursors in the presence of saturating amounts of the other components. The concentration of ribosomes used in the screening assay. (B) Plot of poly(Phe) synthesis as a function of increasing concentrations of *E. coli* EF-Tu (0.023 to 3.0 μM) in the A/T system. The arrow indicates the concentration of EF-Tu used in the screening assay. (C) Determination of the amount of SPA beads needed to quantify the signal from poly(Phe) synthesis in the A/T reactions. The arrow indicates the amount of SPA beads used in the screening reactions.

**RESULTS**

Development and optimization of the A/T assay for screening. An aminocyclation/translation (A/T) system that contained the components required for the translation of poly(U) mRNA—ribosomes, EF-Tu, EF-Ts, EF-G, and PheRS—was developed. All the components of the A/T assay were purified to near-homogeneity as described in Materials and Methods. The coupled A/T reaction was adapted from separate aminocyclation and translation assays (5, 6, 11). The assay was optimized for inhibitor screening in a 96-well microtiter plate format. Ribosomes were initially titrated in the presence of saturating amounts of the other components (Fig. 1A), and 0.1 μM was chosen as the screening concentration that yielded sufficient signal over the background level. At this concentration of ribosomes, all other components of the system were then individually titrated into the system to determine the inflection point of saturation on a titration curve (see Fig. 1B for an example). Concentrations were set just below the saturation points to facilitate the maximum sensitivity to inhibition of each and every component of the system. In the initial screening assays, crude *E. coli* tRNA was used, but secondary-assay mixtures contained purified *E. coli* tRNA^Phe^. A scintillation proximity assay (SPA) was developed for the initial screening assay. The RNA portion of ribosomes was used to localize the ribosome to scintillation beads, enabling the detection of the nascent radiolabeled polyphenylalanine [poly(Phe)] peptide still attached to the ribosome. Yttrium silicate RNA binding beads (Perkin-Elmer) at 200 μg/well gave optimal results. Figure 1C shows the titration of SPA beads in assays to determine the amount of beads required for the capture of the maximum amounts of ribosomes allowing analysis of the activity of the system. The optimal pH for ribosome-bead binding was determined to be 6.2. In the absence of ribosomes, negligible amounts of tRNA charged with [3H]phe- nylalanine or free [3H]phenylalanine bound to the beads.

**Rationale and development of positive controls.** We selected the macrolides spiramycin and tylosin, each containing a 16-member lactone ring and a disaccharide at the C-5 position on the lactone ring, as control antibiotics. Both of these antibiotics inhibited the synthesis of poly(Phe), apparently by binding the lactone ring same with tylosin. The 16-member lactone rings are effective at inhibiting poly(U) synthesis in the A/T assay, with IC_{50} of 0.022 μM and 0.038 μM, respectively (Fig. 2).
Screening of chemical compounds yields primary hits composed of compounds containing 4H-pyridopyrimidine central scaffolds. A chemical-compound library containing 2,100 compounds from Asinex (Moscow, Russia) was tested in a high-throughput format. The screening concentration was 64 μM, and the reactions were carried out as described in Materials and Methods. Nine compounds were observed to inhibit more than 50% of poly(Phe) synthesis. Inhibition of more than 50% of poly(Phe) synthesis defines a hit compound. The ability to inhibit poly(Phe) synthesis was confirmed in triplicate assays for all nine compounds. Structural inspection of the nine compounds revealed that all had the same central scaffold, a tetrahydropyrido[4,3-d]pyrimidin-4-ol (e.g., 4H-pyridopyrimidine) (Fig. 3). In addition, all the structures contained an ortho-pyridine in position 2 of the central scaffold and held all structural variations in a heteroaromatic ring (A-ring) that connected to the N-6 atom via a methylene group. The 2,100-individual-sample library contained numerous chemotypes, of which approximately 113, or 5.4%, were of the 4H-pyridopyrimidine class. This small set of 113 4H-pyridopyrimidines contained an approximately equal distribution of regioisomers (e.g., ortho, meta, para) of the pyridyl ring connected at position 2 of the central scaffold. Other than the nine hit compounds, a subset exhibited activity inhibiting less than 50% of poly(Phe) synthesis, and another subset exhibited no inhibitory activity.

The nine compounds were serially diluted from 200 μM to 0.4 μM, and IC50s were determined. The most potent inhibitors of the A/T reaction were REP321436 and REP321437, with IC50s of 3.7 and 4.9 μM, respectively (Fig. 4). The IC50s of all the test compounds ranged from 3.7 to 14.0 μM (Table 1).

Determination of the target of 4H-pyridopyrimidines. In our system, poly(Phe) synthesis is inhibited by the 4H-pyridopyrimidines. Other than the ribosome, four accessory proteins are required for poly(Phe) synthesis. We used specialty assays to rule out the possibility that the function (or functions) of one of the accessory proteins was inhibited (see below). The 4H-pyridopyrimidines were not observed to inhibit any of the four accessory proteins.

First, the compounds were tested to determine if they inhibited the activity of PheRS. The assay tested the ability of PheRS to attach phenylalanine to its cognate tRNA Phe. Purified E. coli tRNA Phe was used in these assays as described in Materials and Methods. No inhibition of PheRS activity by any of the compounds was observed at compound concentrations as high as 200 μM (data not shown).

Next, the abilities of the 4H-pyridopyrimidines to inhibit the function of EF-Tu were tested. In the absence of EF-Ts and GTP, EF-Tu binds GDP, and the exchange of the bound GDP for free GDP can be monitored. This binding has historically been used to characterize the activities of EF-Tu molecules from various species (33). In these assays, 3H-labeled GDP was used to track the amount of GDP exchanged by EF-Tu and the abilities of the test compounds to inhibit this exchange. We observed no reduction in the exchange of bound and free GDP by EF-Tu in the presence of any of the test compounds at concentrations as high as 200 μM (data not shown). In the presence of EF-Ts, the turnover of GDP binding by EF-Tu is stimulated 5-fold (51). Thus, when EF-Ts is added to the
EF-Tu/GDP exchange assay mixture in the presence of the test compounds, inhibition of the exchange stimulated by EF-Ts may be monitored. The concentration of EF-Tu was reduced from 12 μM to 0.75 μM, and the time for completion of the assay was reduced from 30 min to 30 s. EF-Ts was added to a concentration equal to 1% of the concentration of EF-Tu (0.0075 μM). The stimulatory activity of EF-Ts was not affected by the presence of any of the test compounds at concentrations as high as 400 μM (data not shown).

When EF-Tu, GTP, and aminoacylated tRNA are associated in a ternary complex, EF-Tu protects the acceptor stem of the tRNA and the attached amino acid from nuclease digestion (RNase A). The RNase A assay, as described in Materials and Methods, was used to monitor the ability of EF-Tu to form a ternary complex and protect aminoacylated tRNA (5). E. coli tRNA<sub>Phe</sub> was aminoacylated using [3H]Phe, yielding [3H]Phe-tRNA<sub>Phe</sub>, and the radiolabeled aminoacylated tRNA was subsequently used to test the effects of the test compounds on ternary-complex formation. The compounds were assayed to concentrations as high as 400 μM, and no effect on the ability of EF-Tu to form a ternary complex and protect the bound tRNA was observed (data not shown).

EF-Tu delivers the aminoacylated tRNA to the A-site of the ribosome in the form of a ternary complex. The binding of the cognate tRNA to the A-site stimulates the hydrolysis of GTP by EF-Tu, and the tRNA is subsequently released for sole binding at the A-site. When GTP is replaced with a nonhydrolyzable analog (GDPNP), the ternary complex binds the A-site. When GTP is replaced with a nonhydrolyzable GDP, the ternary complex binds the A-site and the tRNA is subsequently released for sole binding at the A-site. The binding of the ternary complex to the ribosome is stimu-

### TABLE 1. Inhibitory potencies of 4H-pyridopyrimidines against bacterial protein synthesis

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<tr>
<th>Compound</th>
<th>IC₅₀ (μM)</th>
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**4H-pyridopyrimidines specifically inhibit bacterial protein synthesis.** An ideal antibacterial compound would show potent inhibition of its bacterial target but little or no inhibition of the corresponding eukaryotic system. Wheat germ extract assays were used to determine whether the 4H-pyridopyrimidines inhibited protein synthesis in a eukaryotic system. Poly(U) mRNA, yeast tRNA<sub>Phe</sub>, [3H]phenylalanine, and Mg<sup>2+</sup> concentrations were optimized for poly(Phe) synthesis in wheat germ extract assays. The 4H-pyridopyrimidines were tested along with a known inhibitor of protein synthesis, cycloheximide (41). In these assays, cycloheximide inhibited 80% of the pro-

FIG. 4. IC₅₀ determination for protein synthesis inhibitors. The titration of representative 4H-pyridopyrimidines is shown. REP321436 (A) and REP321437 (B) exhibited IC₅₀s of 3.7 and 4.9 μM, respectively, in A/T reactions. The reactions were carried out as described in Materials and Methods, with the concentrations of the compounds ranging from 0.4 to 400 μM. Positive controls were assays in which there were no antibiotics and approximately 1,000 pmol of phenylalanine was synthesized. The curve fits and IC₅₀s were determined using XLfit (version 4.1; IDBS) as part of Microsoft Excel.
tein synthesis at 30 μM. In contrast, none of the test compounds inhibited protein synthesis at concentrations up to 300 μM (Fig. 5). The level of poly(Phe) synthesis in the wheat germ assays is only approximately 10% of that seen in the A/T assays; this is due to the limited number of ribosomes present in the wheat germ lysate.

The compounds were negative in assays carried out to determine if they nonspecifically bind nucleic acid and thereby inhibit mRNA interactions with the ribosome. None of the compounds were shown to interact with nucleic acid nonspecifically (data not shown).

**Microbiological testing of the 4H-pyridopyrimidines.** The 4H-pyridopyrimidines were tested in broth microdilution assays to determine their MICs. Despite the similarity in the biochemical potencies of the compounds, their abilities to inhibit bacterial growth differed widely. None of the test compounds to determine if RNA, DNA, cell wall, lipid, or protein synthesis was inhibited in bacterial cultures. Assays were carried out in cultures containing the E. coli tolC mutant and in cultures of S. aureus. The MMS data for two representative compounds, REP323219 and REP323370, show that the 4H-pyridopyrimidines are specific inhibitors of protein synthesis in the cell (Fig. 6). These molecules are similar to REP321525 except that the fluorine in the A-ring was replaced with chlorine and moved from the meta to the para or ortho position, respectively, relative to the attachment carbon (see Fig. 3). These compounds were tested as we had begun producing modifications of the original hits and focused on testing only the more potent compounds. REP323219 preferentially inhibited protein synthesis in E. coli tolC with an IC50 of 5.6 μg/ml. REP323370 demonstrated a similar MMS profile; measurements from the inhibition plots of the precursor incorporation assays resulted in IC50s of >64 μg/ml for the synthesis of RNA, DNA, the cell wall, and lipids, and an IC50 of 5.8 μg/ml for protein synthesis. The results did not change significantly if the cultures contained E. coli tolC mutants or S. aureus bacteria (data not shown). This shows that the 4H-pyridopyrimidines are broad-spectrum inhibitors and preferentially inhibit protein synthesis in both Gram-negative and Gram-positive bacteria.

**TABLE 2. Antibacterial activities of 4H-pyridopyrimidines against selected pathogens**

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<th>Pathogen</th>
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<td>ND</td>
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*ND, not determined.*
6 represent indirect effects of the inhibition of protein synthesis. Assays with the known protein synthesis inhibitor tylosin gave similar results (data not shown).

Minimum bactericidal concentration (MBC) testing of the 4H-pyridopyrimidines initially indicated that the compounds were bactericidal against *H. influenzae* but only bacteriostatic against *S. pneumoniae*. Time-kill assays with REP321525 at 4× MIC confirmed these results (Fig. 7). A >1,000-fold decrease in viable cell counts within 24 h, which is indicative of bactericidal effects, was achieved upon exposure of *H. influenzae* to REP321525. However, in the time-kill assays, the compound exhibited bacteriostatic effects against both *S. pneumoniae* and *M. catarrhalis*.

The 4H-pyridopyrimidine compounds were also evaluated for their potential to cause nonspecific membrane damage and/or cell lysis. Live/Dead BacLight bacterial viability kits were used to determine whether the test compounds affected the integrity of the bacterial membrane. In *E. coli tolC* mutants, the test compounds did not cause cell lysis at concentrations as high as 128 μg/ml. Similarly, the compounds had no adverse effects on eukaryotic cell membranes, since hemolysis was not observed upon exposure of equine erythrocytes to this class of compounds at 128 μg/ml.

### DISCUSSION

**Biochemical screening and target identification of bacterial protein synthesis inhibitors.** The results presented here show that a coupled aminoacylation/translation (A/T) system constructed using purified components is functional in poly(Phe) synthesis and can be used to screen for compounds that inhibit protein synthesis in bacteria in a high-throughput format. We have used this system to screen a small-molecule compound library containing 2,100 compounds, and we identified a series of 4H-pyridopyrimidine analogs that are bacterial protein synthesis inhibitors. All of the compounds in the library had molecular masses of approximately 500 Da. Through the use of a set of assays to determine the effects of the 4H-pyridopyrimidines on the activities of the ribosomal ligands, we have determined that the likely target of the inhibitors is the ribosome. The activities of EF-Tu, EF-Ts, EF-G, and PheRS showed no sign of a reduction when assayed in the presence of test compounds, whereas in A/T assay mixtures containing ribosomes, protein synthesis was reduced in a dose-dependent manner.

Using macromolecular synthesis assays, the mode of action was confirmed to be inhibition of protein synthesis. In contrast, the 4H-pyridopyrimidines had little or no effect on DNA, RNA, cell wall, or lipid production in bacteria, and the syn-
thesis of these macromolecules was affected only indirectly due to blocking of the production of proteins essential for their synthesis. In addition, the 4H-pyridopyrimidines did not block protein synthesis in eukaryotic cells. Inhibition of protein synthesis in bacteria as a mode of action with the ribosome as the target and specificity for bacteria makes the 4H-pyridopyrimidines candidates for drug development against pathogenic bacterial infections.

4H-pyridopyrimidines are active in microbiological assays.

The compounds identified in the A/T screen were subjected to microbiological assays in which they inhibited bacterial growth. Initial assays indicated that the more potent compounds had activity against S. aureus, S. pneumoniae, and permeabilized or efflux-compromised E. coli. Secondary assays confirmed the broad-spectrum activity of this compound series against both Gram-positive and Gram-negative bacteria (Table 2). In particular, the 4H-pyridopyrimidines had good activity against all the respiratory pathogens tested, including S. aureus, S. pneumoniae, H. influenzae, and M. catarrhalis. In time-kill assays, the compounds were shown to have bacteriostatic activity against S. pneumoniae and M. catarrhalis but were bactericidal against H. influenzae. In general, inhibitors of protein biosynthesis tend to exhibit a static effect on the growth of bacteria; however, some protein synthesis inhibitors, such as certain aminoglycosides, do exhibit bactericidal effects (20). The bacteriostatic/bactericidal effects of 4H-pyridopyrimidines appear to be bacterial strain specific.

Antibiotics targeting the ribosome and mechanisms of action.

Previous work indicates that chloramphenicol and certain macrolides fail to inhibit poly(Phe) synthesis in cell extracts (25, 32, 34). We initially tested azithromycin and chloramphenicol in the A/T system, and a <5% decrease in poly(Phe) synthesis was detected. More-recent evidence suggested that macrolides containing a 14-member lactone ring structure inefficiently inhibited the peptidyl transferase reaction, and certain macrolides containing a 16-member lactone ring but only a single sugar moiety at C-5 also failed to inhibit peptidyl transfer completely (36). These macrolides apparently bind in the peptide exit tunnel adjacent to the PT center, and in normal protein synthesis, they allow 6 to 8 amino acids to be polymerized, as has been previously observed for both tylosin and spiramycin (4). This may be a characteristic of the synthesis of pure poly(Phe), which is atypically hydrophobic compared to a natural protein sequence (32, 34). We have not yet determined the exact mode of action of the 4H-pyridopyrimidines on the ribosome, but the levels of residual activity observed in poly(Phe) synthesis were similar to the levels observed with tylosin and spiramycin. One possibility is that the 4H-pyridopyrimidine may be binding in or near the exit tunnel for the polypeptide or near the PT center, allowing several amino acids to be polymerized before synthesis is completely blocked. Additional structure/function studies will be needed to elucidate the mechanism of action.

Other compounds in this class have a different mode of action. Concurrent with the work described here, in an attempt to elucidate inhibitors of bacterial growth, Miller et al. at Pfizer (Ann Arbor, MI) screened a 1.6 million-compound library against E. coli (tolC imp) in whole-cell assays (27). A series of compounds with a pyridopyrimidine scaffold was identified and was shown to inhibit the activity of bacterial bacteriophage. By use of macromolecular synthesis assays, the mode of action was confirmed to be inhibition of fatty acid synthesis. In contrast, the 4H-pyridopyrimidines described here had no effect on the synthesis of lipids (Fig. 6B). Conversely, the Pfizer compounds had no effect on protein synthesis. There are subtle differences in the electronic configuration of the pyridine ring structure within the central scaffold between the two series of compounds. The Pfizer compounds also lack the pyridine ring attached to the central scaffold that is present in the 4H-pyridopyrimidine series, and even though the central scaffolds of the two compound series are structurally similar, the site of action is completely different.

The A/T system as a screening platform.

Initially, nine compounds from a selected low-molecular-mass chemical-compound library were identified as inhibitory in the A/T system. The results shown here demonstrate that a reconstituted protein synthesis system composed entirely of purified components can be an effective screening system. The resulting hits can be characterized through a series of assays to identify the target component. This system can be applied to the screening of larger chemical-compound libraries and offers significant advantages over reactions carried out using crude extracts. The A/T system has the potential to be adapted to a 384-well microtiter plate format with further optimization of enzymatic and detection components, allowing higher-throughput screening. In addition, this system can be used to aid in the identification of targets of lead compounds identified in whole-cell screens with a mode of action of inhibiting protein synthesis.

The A/T system has obvious limitations in that only inhibitors of poly(Phe) synthesis will be detected. In the translation of a natural mRNA, 19 additional aminoacyl-tRNA synthetases and several additional initiation and termination factors are required. To detect inhibitors of all components of protein synthesis, a more complete system will be required.

To proceed in the development of the 4H-pyridopyrimidines as drug candidates requires in-depth development. During follow-up studies, the 4H-pyridopyrimidines were subjected to further medicinal chemistry optimization to maximize antibacterial activity, and during the course of this work, more than 100 additional analogs have been synthesized. The IC50s for these optimized compounds have been reduced in the A/T...
protein synthesis system to the low nanomolar range, and the MICs for *H. influenzae*, *S. pneumoniae*, *M. catarrhals*, and *S. aureus* have been reduced to 1, 2, <0.12, and 4 μg/ml, respectively. These studies will be discussed elsewhere.

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REFERENCES


