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Accessibility
Identification of DNA primase inhibitors via a combined fragment-based and virtual screening

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The structural differences between bacterial and human primases render the former an excellent target for drug design. Here we describe a technique for selecting small molecule inhibitors of the activity of T7 DNA primase, an ideal model for bacterial primases due to their common structural and functional features. Using NMR screening, fragment molecules that bind T7 primase were identified and then exploited in virtual filtration to select larger molecules from the ZINC database. The molecules were docked to the primase active site using the available primase crystal structure and ranked based on their predicted binding energies to identify the best candidates for functional and structural investigations. Biochemical assays revealed that some of the molecules inhibit T7 primase-dependent DNA replication. The binding mechanism was delineated via NMR spectroscopy. Our approach, which combines fragment based and virtual screening, is rapid and cost effective and can be applied to other targets.

The complex process of identifying antibacterial compounds begins with the selection of potential targets, which must be essential, selective over human homologues, susceptible to drugs, and with a low propensity to develop rapid resistance1. Although bacteria possess approximately 200 essential gene products, only a limited number of these have been exploited as drug targets2. DNA replication, which qualifies as a novel drug target, is performed by the replisome, a multi enzyme complex that synthesizes DNA continuously on its leading strand and discontinuously on its lagging strand3,4. DNA primase, an essential component of the DNA replication machinery of every living cell5, synthesizes short RNA primers that are used by DNA polymerase to form the “Okazaki fragments” on the lagging DNA strand. The inhibition of primase, therefore, will halt DNA replication and, as a result, cell proliferation.

Prokaryotic primases (among which is the primase domain of the multifunctional gene 4 protein of bacteriophage T7, the model used in our study) share a conserved primary sequence (Fig. 1a) and are structurally highly similar (Fig. 1b)5. In contrast, the profound differences between human and bacterial DNA primases (Fig. 1c) render the latter a selective target for drug design. Specifically, human primase has four subunits6 (Fig. 1c, right), while bacterial DnaG usually functions together with the hexameric ring of DnaB helicase (Fig. 1c, left). In addition, sequence homology between the mammalian and bacterial primases is very low5. Finally, DnaG possesses an active site for binding nucleotides and a DNA binding module, indicating that bacterial primase is a target for drugs. Despite its high therapeutic potential, however, no clinical candidate inhibitors of DnaG primase have emerged to date.

Historically, the screening process for potential ligands has relied heavily on high throughput screening (HTS). The low effectiveness of HTS in identifying new antibacterial agents7, however, led to the emergence of fragment-based screening as a viable alternative route for hit discovery in infectious disease research. Screening of small molecules, whether by fragment screening or HTS, can target key biochemical process or binding to an essential cellular component. Fragment-based screening monitors the binding of smaller molecules from fragment libraries8, where the small sizes of the molecules constituting a typical fragment library increases the chances of binding but the strength of that interaction is weak9,10. Another potential disadvantage for fragment-based screening is the low selectivity of the resulting hits11. Although such low affinity-low selective-low weight hits were not believed to indicate the presence of a potentially viable clinical candidate, molecules found by using

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fragment-based screening are emerging in the late stages of clinical trials8. To detect the weak binding affinities (KD ~ μM - mM) on which fragment-based screening relies, saturation transfer difference (STD) spectroscopy is used12.

The weak catalytic activity of DNA primase renders the adaptation of a functional assay to HTS a formidable challenge. Here we propose a novel, hybrid method for developing small molecule inhibitors for T7 primase to circumvent the drawbacks of HTS (Fig. 2). Based on the ‘rational design’ philosophy of lead development, our method exploits NMR to identify binders from libraries of fragment molecules. We then use computational methods to construct larger molecules with improved binding/inhibition properties. We show that the use of fragment based virtual screening (FBVS, Fig. 2) can yield potent inhibitors, reduce costs, and provide more advanced information about lead binding properties prior to the medicinal chemistry phase of drug optimization.

Results
Bacterial DnaG primase synthesizes RNA primers that are used by DNA polymerase in lagging strand synthesis during DNA replication. Owing to the stark differences between the human and bacterial primases, DnaG primase has long been a target for drug discovery. Here we describe the development of small molecule inhibitors of DnaG using the structurally similar primase domain of the bacteriophage T7 gene 4 protein (Fig. 1b) as a model for the bacterial primase.
To find compounds that do not bind at high affinity but that have potential to become successful leads, we developed a platform for lead discovery comprising several complementary steps: (1) fragment based screening by STD spectroscopy of a small molecule library (Ro3), (2) hit optimization by virtual screening and the generation of a new set of drug-like compounds (Ro5), each of which contains a small molecule found in step 1, (3) docking of the drug-like compounds to the active site of T7 primase and selection of the compounds that will undergo functional and structural assays with the target T7 primase, and (4) further development of lead compounds.

**Fragment Based Virtual Screening (FBVS) – A combined screening approach.** Growing a fragment molecule into a larger molecule that possesses drug like properties is the bottleneck in fragment-based drug design approaches. Using the Maybridge Ro3 fragment library containing 1000 fragments, we prepared 100 NMR samples, each containing a mixture of 10 fragments and 50 μM T7 primase. We used 10 fragments per each NMR sample to significantly minimize NMR time. The 10 fragments were chosen to show minimal overlap of their 1H chemical shift to allow easy identification of the shifts. The 1D saturation transfer difference spectra of these samples were measured, and fragments that exhibited saturation transfer (evident by a decrease in the peak intensity) were identified. Hits were ranked based on the number of peaks in the 1H-NMR spectra with decreased intensities. The small molecule fragments indole and 2H-chromene-3-carbothioamide (Fig. 3) exhibited the strongest binding that, correspondingly, was reflected in the largest decreases in NMR peak intensities (considering the total change in intensity and also the number of affected peaks). Hits were validated by measuring the [15N,1H] HSQC spectra of 15N,D-T7 primase and then evaluating the chemical shift perturbations of the backbone amide resonances upon the addition of the small fragment molecules (Fig. 3). This validation also ensures that the change in peak intensities in the 1D spectra was not a result of additive effects of several fragments.

**Lead optimization and candidate selection.** Hits from fragment-based screening are commonly optimized by creating larger compounds with better binding properties, a time and resource intensive step that usually involves medicinal chemistry.

To eliminate the need for a medicinal chemistry phase during the early steps of lead optimization, we used virtual screening where the structure of the small molecule fragment binder found by STD spectroscopy was used as a constraint for the next search procedure. We searched the rapidly growing ZINC database, which contains the structures of over 100 million compounds, for those with at least 70% similarity to the fragment molecules identified by STD spectroscopy. The two fragment molecules ranked highest by the fragment based screening – indole and 2H-chromene-3-carbothioamide – were used for this step of virtual filtration. The database search yielded approximately 3000 compounds per scaffold. We then used the docking software AutoDock to dock these compounds into the active site of T7 primase using its available crystal structure (pdb code 1nui). The compounds were then ranked based on their ΔG predicted binding energy values. The highest-ranking compounds that
Small molecules that inhibit the concerted activity of primase and DNA polymerase. To evaluate the ability of the small molecule candidates to inhibit the primase-dependent replication of the lagging strand of bacteriophage DNA, we ran an overall assay that involved the concerted activity of DNA polymerase (gene product 5 and \( E. coli \) thioredoxin, gp5/trx) and the helicase-primase (gene product 4, gp4, full-length). In bacteriophage T7, lagging strand DNA synthesis involves interactions between gp5/trx and gp4. Full-length gp4 is required for the synthesis of oligoribonucleotides to initiate the synthesis of Okazaki fragments. To examine the effect of 350 \( \mu M \) of each small molecule on the synthesis of primers and their transfer to gp5/trx, we used M13 ssDNA for the synthesis of oligoribonucleotides by the primase and their extension by gp5/trx (Fig. 4b, inset). To initiate DNA synthesis, the primase must first synthesize tetraribonucleotides on the DNA and then transfer them to gp5/trx. In addition to the four dNTPs, ATP and CTP were also provided, and the primers that were synthesized included pppACCC, pppACAC, and pppACCA. Inhibition of the primase activity of gp4 halts RNA primer formation process, thereby preventing subsequent DNA polymerase activity. Figure 4 shows that primase dependent DNA synthesis decreases by up to four-fold with the addition of five small molecules, including (2E)-3-(6-chloro-2H-chromen-3-yl)acrylic acid (compound 1), 9-Nitro-7,12-dihydroindolo-[3,2-d]
benzazepin-6(5)-one (compound 12), 3-[2-(ethoxycarbonyl)-5-nitro-1H-indol-3-yl]propanoic acid (compound 13), N-(1,3-benzodioxol-5-yl)-7-nitro-1H-indole-2-carboxamide (compound 15), and 7-nitro-1H-indole-2-carboxylic acid (compound 17), whose chemical structures are presented in Fig. 4a.

Small molecules that inhibit primase activity. To catalyze the synthesis of short RNA primers, T7 primase requires DNA with a primase recognition site, ATP and CTP, and a buffer containing divalent metal ions. The effect of each of the five small molecules found using FBVS approach as described above on RNA primer synthesis by the T7 primase was examined. The primase domain catalyzed the synthesis of the diribonucleotide pppAC on a DNA template containing the 5',GTC,3' primase recognition site. We examined diribonucleotide synthesis in the presence of each of the five small molecules found to inhibit the concerted activity of primase and polymerase (Fig. 4b). The reaction conditions involved incubating the T7 primase with an oligoribonucleotide containing a primase recognition sequence, \([\alpha-32P]\)-CTP and ATP, and adding each small molecule in steadily increasing amounts. The radioactively labeled oligoribonucleotides were separated on a denaturing polyacrylamide gel, and radioactivity was measured on an autoradiogram. The results clearly show that inhibition of the specific activity of the primase increased with increasing amounts of the small molecules (Fig. 4c). Curve fitting was performed using nonlinear four parameter logistics to explore features of small molecule binding to the...
enzyme. Values for IC₅₀ and Hill coefficient were extracted (Fig. 4c, right bottom). Data for compounds 1 and 13 were not sufficient for the analysis therefore IC₅₀ values and Hill coefficients were excluded. Overall, IC₅₀ for those molecules were in the sub-millimolar range. Hill coefficient for compound 12 present highest value presumably due to stronger binding of the molecule to the enzyme.

Structural analysis of inhibitor-primase interactions. To characterize the binding of the compounds identified using FBVS, we measured the [¹⁵N, ¹H]-TROSY-HSQC spectra of ¹⁵N- and D-labeled T7 primase in the presence of the selected lead compounds (Fig. 5). After the addition of compounds 1, 13, and 17 to T7 primase, we observed significant chemical shift perturbations compared to those obtained for the free T7 primase. Relative to the initial scaffold alone, those shifts were more pronounced, which indicates stronger binding interactions. In addition, chemical shift perturbations upon the addition of DNA and ATP/CTP to the T7 primase domain confirmed that all three compounds bind to the active site. Moreover, the presence of several cross peaks among the three compounds indicates that they exploit a similar binding mechanism. Compounds 12 and 15, whose polar dissolution values were the lowest from among the five inhibitors, were not able to achieve the concentrations needed for the protein-NMR experiments.
Figure 5. Binding-site of small-molecule inhibitors on T7 DNA primase. (a) Amino acid sequence chemical shift assignments indicate that small molecule binding occurs in the proximity of the main cleft of T7 DNA primase (binding site indicated in green). (b) Right: Two-dimensional $^1$H-$^{15}$N HSQC spectrum of T7 DNA primase alone (black spots) and in the presence of each small molecule inhibitor (blue, compound 1; purple, compound 13; green, compound 17). The HSQC spectrum of primase changes in the wake of small molecule binding. Similar peaks change upon titration of DNA (GGTCA) or ATP and CTP in the presence of magnesium added in a ratio molar concentration. Left: Amino acid residues that mediate the binding of each small molecule inhibitor are indicated in green.
The fragment based screening approach proposed here enabled us to rationally design drug like inhibitors (that contain small fragments) to the primase. However, in contrast to conventional fragment-based screening, which requires a subsequent medicinal chemistry step to grow the hit molecules, FBVS eliminates the need for that step early on and increases the chances of successfully identifying larger molecules containing the fragment hits. Insofar as the inhibition of DNA primase will stop bacterial DNA replication and prevent infection, the primase inhibitor will be at the forefront of a new class of anti-bacterial agents.

To unravel the binding site of T7 primase inhibitors, we assigned the resonances to the T7 primase backbone (see Methods). With the exception of the zinc-binding domain, 70% of the chemical shifts of the T7 primase have been assigned to their corresponding residues (unpublished data). Assignment of the NMR peaks to the protein residues enabled us to identify the amino acid residues situated in the proximity of the active site that mediate small molecule inhibitor binding (Fig. 5a,b). Protein stability was severely impaired if the six amino acid residues at the binding site – Ala(80), Ser(87), Glu(89), Val(101), Met(105), Tyr(106) – are substituted altogether with Ala (except Ala(80), which was substituted with Gly).

Figure 6a shows the amino acid residues that mediate the binding of each of the three small molecule inhibitors [(2E)-3-(6-chloro-2H-chromen-3-yl)acrylic acid (compound 1); 3-[(2-(ethoxycarbonyl)-5-nitro-1H-indol-3-yl)propanoic acid (compound 13); 7-nitro-1H-indole-2-carboxylic acid (compound 17)]. Indeed, for each of the three compounds, the mechanism of binding to the active site cleft is similar but not identical, and the amino acids involved overlap slightly, such that Val(101), Met(105), and Tyr(106) mediate the binding of all three compounds (Fig. 6a). All three of the small molecules bind to the primase active site and are expected to interfere with the binding to the substrate (ribonucleotide tri-phosphates) or to the DNA template. Indeed, substituting two of the amino acid residues that mediate the binding of all three compounds [Val(101) and Met(105)] with Ala inactivated the primase (Fig. 6), an outcome that is indicative of the central roles these two amino acid residues play in primer synthesis. More importantly, this specific binding location represents a potential route to prevent bacteria from evolving into a resistant strain: any adaptive mutation driven by inhibitor binding would completely disrupt primase activity, killing the bacteria as a result.

Discussion

Functional assays for primase generate weak readout signals, and therefore, are not easily adapted to high throughput screening. The T7 primase is an ideal model to study bacterial primases because (1) it shares high structural similarity with bacterial primases, (2) it is highly expressed, (3) it has been extensively studied, and (4) its crystal structure is known. We assigned NMR HSQC perturbations to the amino acid sequences of the primase, which enabled us to identify the amino acid residues that mediate small molecule inhibitor binding. This knowledge will be invaluable for further optimization and structure activity relationship studies, which have the potential to lead to the development of new antibacterial drugs.

In this study, we introduced FBVS, a novel analytical technique that combines a spectroscopic assay (STD spectroscopy) with an in-silico approach. Its computational component comprised the virtual filtration of a small molecule database followed by the docking of the potential small molecule inhibitors to the crystal structure of T7 primase. Not only did this approach allow us to find several lead compounds for T7 primase inhibition, it also showed a high hit rate: of the 16 compounds ordered for testing, about half showed primase inhibition activity. This approach, which is not only fast but also cost effective, is a promising method for identifying the inhibitors of protein targets that are not amenable to high throughput screening.

Figure 6. Essentiality of amino acids that mediate the binding of small molecule inhibitors. (a) Amino acid residues that mediate small molecule inhibitors at the active site of T7 DNA primase. The three T7 primase inhibitors share the same binding site and are mediated via similar binding mechanisms. Solvent accessible amino acid residues were calculated using Naccess (http://www.bioinf.manchester.ac.uk/naccess/) and are marked in asterisk. (b) Substitutions of the central amino acids that were shown to mediate the binding of all the tested inhibitors and are solvent accessible (i.e., Val101 and Met105 were replaced with Ala) disrupted protein activity. The faint signal of pppAC remained for the reaction of the double mutant T7 DNA primase may be due to erroneous loading of the sample into the gel. Reaction conditions were as in Fig. 4 (except that the protein concentrations were 0.2, 0.6, 1.9, 3.8 μM, respectively). (c) Quantification of di-ribonucleotide synthesis by T7 DNA primase. The bands in the gels presented in b were analyzed using autoradiography.

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The fragment based screening approach proposed here enabled us to rationally design drug like inhibitors (that contain small fragments) to the primase. However, in contrast to conventional fragment-based screening, which requires a subsequent medicinal chemistry step to grow the hit molecules, FBVS eliminates the need for that step early on and increases the chances of successfully identifying larger molecules containing the fragment hits.

Insofar as the inhibition of DNA primase will stop bacterial DNA replication and prevent infection, the primase domain of phage T7 gene 4 product represents an important drug target. We therefore expect that a T7 DNA primase inhibitor will be at the forefront of a new class of anti-bacterial agents.
Methods

Protein expression and purification. Chemicals were from Sigma. ATP and CTP were from Roche Molecular Biochemicals. dNTPs and ddNTPs were from New England Biolabs. dNTPs and ddNTPs were purchased from Boehringer Mannheim. M13 ssDNA was prepared as described previously24. [γ-32P] dATP (80 Ci/mmol), [α-32P] CTP, and dTTP (80 Ci/mmol) were from Perkin Elmer.

Protein expression and purification. Proteins and Reagents. All chemical reagents were of molecular biology grade (Sigma); ATP and CTP (Roche Molecular Biochemicals). dNTPs and ddNTPs were purchased from Boehringer Mannheim. Premade gels (10–20% linear gradients) used for SDS–PAGE and Precision Plus Protein prestained standards were purchased from BioRad (Hercules, CA). T7 primase domain (residue: 1–271) was over-produced and purified using metal free buffers as previously described17,21. T7 gp5 and E. Coli trx were overproduced and purified as described18. Gp4 was overproduced and purified as described19. M13 ssDNA was prepared as described previously20. [α-32P]–CTP (800 Ci/mmol) was purchased from Perkin Elmer.

Virtual screening. The hits determined by NMR were used to identify compounds with at least 70% similarity from the ZINC database22. Virtual screening was performed to identify molecules that could bind in the active site of T7 DNA primase and inhibit DNA replication. Docking of all compounds downloaded from the ZINC database was performed using AutoDock4.2.16. PDB files of the receptor (T7 primase) and the ligands (compounds) were prepared prior to docking as described in the tutorial “Using AutoDock for Virtual Screening” (http://autodock.scripps.edu/faqs-help/tutorial/using-autodock-for-virtual-screening/UsingAutoDockforVirtualScreening_v7.pdf). The search grid was centered in the active site of T7 primase with a grid spacing of 0.37 Å and 110 × 108 × 126 points. The default parameters were used except for the following modified parameters: ga_num_evals = 1750000, ga_pop_size = 150, and ga_run = 100.

Primase dependent DNA synthesis. RNA primers made by gp4A were extended by gp5/trx. The reaction mixture contained 10 nM M13 ssDNA, 0.3 mM dNTPs, 0.1 μCi [α-32P] dCTP, 20 mM gp5/trx, 200 mM monomeric gp4A, and 350 μM of each compound. The reaction was incubated for 45 min at 37 °C. Reaction was terminated and amount of DNA synthesis was determined as described in DNA polymerase assay.

DNA polymerase assay. DNA polymerase activity was measured in a reaction containing 5 mM gp5/trx, 20 nM M13 ssDNA annealed to a 24mer primer, 40 mM Tris–HCl (pH 7.5), 10 mM MgCl2, 10 mM DTT, 50 mM potassium glutamate, 0.25 mM dTTP, dGTP, dCTP, [α-32P] dATP (5 cpm/pmol) and 500 μM of each compound. The reaction was incubated at 37 °C for 20 min and terminated by the addition of EDTA to a final concentration of 40 mM. Aliquots of the reaction were spotted on DE–81 filters (Whatman), washed three times with excess of 0.3 M ammonium formate (pH 8.0), and the radioactivity retained on the filters measured.

Oligoribonucleotide Synthesis Assay. Oligoribonucleotides were synthesized by DNA primase, product was measured as described22,23 in reactions containing various concentrations (1.1, 3.3, and 10 μM) of gene 4 primase domain. Standard 10 μL reaction contained 5 μM of DNA template (5′-GGGTCA10′-3′), 200 μM ATP, 200 μM [α-32P]–CTP, and primase domain in a buffer containing 40 mM Tris–HCl (pH 7.5), 10 mM MnCl2, 10 mM DTT, and 50 mM potassium glutamate. After incubation at room temperature for 20 minutes, the reaction was terminated by adding an equal volume of sequencing buffer containing 98% formamide, 0.1% bromophenolblue, and 20 mM EDTA. The samples were loaded onto 25% polyacrylamide sequencing gel containing 3 M urea and visualized using autoradiography.

Sample preparation. For the expression of unlabeled proteins, the primase domain of the bacteriophage T7 gene 4 product (1–271) was expressed in E. coli B21(DE3) containing pETg4A as reported previously21. For the expression of perdeuterated proteins, the primase domain of the bacteriophage T7 gene 4 product (1–271) was expressed in E. coli BL21(DE3) containing pETg4A, and 350 μM of gene 4 primase domain. Standard 10 μL reaction contained 5 μM of DNA template (5′-GGGTCA10′-3′), 200 μM ATP, 200 μM [α-32P]–CTP, and primase domain in a buffer containing 40 mM Tris–HCl (pH 7.5), 10 mM MnCl2, 10 mM DTT, and 50 mM potassium glutamate. After incubation at room temperature for 20 minutes, the reaction was terminated by adding an equal volume of sequencing buffer containing 98% formamide, 0.1% bromophenolblue, and 20 mM EDTA. The samples were loaded onto 25% polyacrylamide sequencing gel containing 3 M urea and visualized using autoradiography.

NMR experiments –Fragment Based Screening. T7 primase was screened against the Maybridge Ro3 Diversity Fragment Library containing 1000 compounds using STD spectroscopy. Samples for 1D ligand-observed NMR studies contained 5 μM T7 primase in 50 mM phosphate buffer in D2O and 10 compounds, resulting in a total of 100 samples. NMR spectra were recorded on a Bruker Avance 500 MHz equipped with a TXO cryoprobe with Z gradient and a NMR-CASE sample changer at 298 K. Ligand binding was probed
using a saturation transfer difference pulse program^{24–26}. Saturation was achieved using on-resonance irradiation at 0 ppm with a train of Wurst pulses for a total saturation time of 1.5 s. Off-resonance irradiation was centered at 40 ppm. Spectra were acquired with a sweep width of 8012.8 Hz, 8192 data points and 224 scans. Active compounds were identified by comparing the STD spectra with the corresponding reference spectra using the MestreNova software. Any compound with reduced peak intensities was considered a hit and ranked according to the percentage of intensity decrease and the number of affected peaks per compound.

[^H,^15N]-TROSY HSQC titration spectra of ^15N,^13C T7 DNA primase with selected fragment molecules and compounds 1, 12, 13, 15, 17, ATP, and DNA were recorded at 25 °C on a Bruker DMX 800 MHz spectrometer equipped with TXI cryoprobes with Z gradient. Data were processed and analyzed using NMRPipe^{27} and NMRView^{28}.

### NMR experiments – Backbone resonance assignments.
Spectra were acquired at 298.15 K with 700μM protein samples in 25 mM KH2PO4/K2HPO4, pH = 7.2, 150 mM NaCl, 1 mM DTT. Traditional TROSY-based backbone triple resonance experiments (HNCA/HNCOCA, HNCO/HNCACO, HNCA) were conducted on a ^15N,^13C-perdeuterated sample to assign the backbone chemical shifts. The spectra were recorded on a Varian 600 MHz spectrometer equipped with cryogenically cooled probe. Non Uniform Sampling was used in all the triple resonance experiments where 12% of the indirect dimension grid was sampled and the spectrum was reconstructed using hmsIST^{29} and NMRPipe^{30}. The resulting spectra were visualized and analyzed using CARA^{31}.

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B.A., S.I., S.R.A., H.A., G.W. and C.C.R. designed research, performed the studies and analyzed the data. B.A. wrote the paper.

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