Title
An ultrasensitive high throughput screen for DNA methyltransferase 1-targeted molecular probes.

Permalink
https://escholarship.org/uc/item/2665z439

Journal
PloS one, 8(11)

ISSN
1932-6203

Authors
Fagan, Rebecca L
Wu, Meng
Chédin, Frédéric
et al.

Publication Date
2013

DOI
10.1371/journal.pone.0078752

Peer reviewed
An Ultrasensitive High Throughput Screen for DNA Methyltransferase 1-Targeted Molecular Probes

Rebecca L. Fagan1, Meng Wu1, Frédéric Chédin2, Charles Brenner1*

1 Department of Biochemistry, Carver College of Medicine, University of Iowa, Iowa City, Iowa, United States of America, 2 Department of Molecular and Cellular Biology and Genome Center, University of California Davis, Davis, California, United States of America

Abstract

DNA methyltransferase 1 (DNMT1) is the enzyme most responsible for epigenetic modification of human DNA and the intended target of approved cancer drugs such as 5-aza-cytidine and 5-aza-2'-deoxycytidine. 5-aza nucleosides have complex mechanisms of action that require incorporation into DNA, and covalent trapping and proteolysis of DNMT isozymes. Direct DNMT inhibitors are needed to refine understanding of the role of specific DNMT isozymes in cancer etiology and, potentially, to improve cancer prevention and treatment. Here, we developed a high throughput pipeline for identification of direct DNMT1 inhibitors. The components of this screen include an activated form of DNMT1, a restriction enzyme-coupled fluorogenic assay performed in 384 well plates with a z-factor of 0.66, a counter screen against the restriction enzyme, a screen to eliminate DNA intercalators, and a differential scanning fluorimetry assay to validate direct binders. Using the Microsource Spectrum collection of 2320 compounds, this screen identified nine compounds with dose responses ranging from 300 nM to 11 μM, representing at least two different pharmacophores with DNMT1 inhibitory activity. Seven of nine inhibitors identified exhibited two to four-fold selectivity for DNMT1 versus DNMT3A.

Citation: Fagan RL, Wu M, Chédin F, Brenner C (2013) An Ultrasensitive High Throughput Screen for DNA Methyltransferase 1-Targeted Molecular Probes. PLoS ONE 8(11): e78752. doi:10.1371/journal.pone.0078752

Editor: Manfred Jung, Albert-Ludwigs-University, Germany

Received June 28, 2013; Accepted September 20, 2013; Published November 13, 2013

Copyright: © 2013 Fagan et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: Work was supported by contract HHSN26120043300C and grant R01CA75954 from the National Cancer Institute and a generous gift from the Roy J. Carver Foundation to CB, grant GM094299 from the National Institutes of Health to FC and an American Cancer Society postdoctoral fellowship to RLF. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: charles-brenner@uiowa.edu

Introduction

In eukaryotes, the most common DNA modification is methylation of the 5 carbon of cytosines, predominately in CpG dinucleotides. Methylation patterns are established and maintained by a family of enzymes known as DNA methyltransferases (DNMTs). De novo methyltransferases, DNMT3A and DNMT3B, establish methylation patterns during germ cell and embryonic development. These proteins are aided by DNMT3L, a catalytically inactive isoform that forms complexes with DNMT3A and DNMT3B [1]. Methylation patterns are primarily maintained by DNMT1, which is the most abundant DNMT and possesses specificity for methylation of hemimethylated DNA [2,3]. DNA methylation is an important epigenetic mark associated with gene repression that plays a critical role in development and differentiation, genome stability, genomic imprinting, X-chromosome inactivation and silencing of retrotransposons [4]. Aberrant DNA methylation has been linked to several diseases including schizophrenia [5], Rett Syndrome [6], autoimmune diseases [6–8], hereditary sensory neuropathy, dementia and hearing loss [9], and cancer [10,11]. In malignancies, normal methylation patterns are disrupted such that global cytosine DNA methylation is reduced, while the regulatory regions of many tumor suppression genes are hypermethylated, resulting in gene silencing [12]. Though genetic changes associated with cancer cannot be corrected, epigenetic changes, such as DNA methylation, are dynamic and amenable to reversal. Epigenetic reprogramming, accomplished by pharmacological targeting of DNMTs, could be expected to result in restoration of a more differentiated and less proliferative state, and regression to a lower degree of drug resistance [13].

The link between the DNMT isozyme DNMT1 and cancer initiation and progression is well established. DNMT1 activity is increased in a variety of malignancies. Several common oncogenic pathways result in the overexpression of DNMT1, either via transcriptional or post-translational mechanisms [14–17] and targeting the DNMT1 isozyme for cancer therapy has been validated genetically. For example, lowering the level of DNMT1 with a Dnmt1 null over Dnmt1 reduced activity genotype protects against tumor formation in Apo−/− mice [18]. In addition, knocking down Dnmt1 with antisense oligonucleotides inhibits neoplasia in cell culture and in mouse tumor models [19,20]. Though genetic experiments can easily target specific DNMT isozymes, this has not been accomplished by pharmacological agents. Discovery of DNMT1 isozyme specific inhibitors could be of great importance as DNMT3A is inactivated in a high proportion of malignancies such as acute myeloid leukemia [21].

Two distinct classes of demethylating agents have been reported. Nucleoside inhibitors such as 5-aza-cytidine and 5-aza-2’-deoxycytidine are FDA-approved prodrugs for treatment of myelodysplastic syndrome [22]. However, these compounds have complicated mechanisms of action that require their incorporation into DNA. Once incorporated, 5-aza nucleotides act as suicide inhibitors, which trap DNMT isozymes in covalent DNA-protein complexes that are cleared by proteolysis and DNA repair, which contributes to the mechanism of action. 5-aza nucleosides are
incorporated nonspecifically into the genome, *i.e.*, not specifically at CpG dinucleotides. Because 5-aza-cytidine metabolism includes formation of 5-aza-CTP and incorporation into RNA, significant toxicity can occur [22,23]. The second class of reported demethylating agents are non-nucleoside inhibitors. This class contains compounds of broad chemical diversity, many of which have other known targets [24]. These agents include compounds that directly inhibit all DNMT isoforms, such as the SAM-competitive inhibitor SGI-1027 [25], and other compounds lacking in direct experimental evidence for a mechanism of inhibition. The cytoxicity and lack of specificity of known DNA demethylating agents suggests a need for new DNMT inhibitors. Thus, we set out to discover novel direct inhibitors of DNMT1 enzyme activity.

Two factors that have delayed discovery of DNMT1 enzyme inhibitors are the intrinsic low activity of the enzyme and the laborious nature of traditional DNA methylation assays utilized to examine DNMT activity [26]. We recently solved both problems by discovering that the replication foci targeting sequence (RFTS) domain is an intrinsic, DNA-competitive inhibitor of DNMT1 enzyme activity and by employing a fluorogenic assay with outstanding signal-to-noise [27]. Two other high throughput screening (HTS)-compatible assays have been recently reported [28,29]. Both assays were used to screen for inhibitors of the de novo methyltransferase DNMT3A. The scintillation proximity assay was also used to screen DNMT3A hits against DNMT1 [29]. Here, we report optimization of an endonuclease-coupled DNMT1 assay to screen a 2320 compound library for small molecules that inhibit DNMT1 enzyme activity. Following validation of initial HTS hits, candidate inhibitors were screened for direct binding of DNMT1 in the absence of substrates using differential scanning fluorimetry (DSF). The pipeline described here resulted in discovery of nine previously unreported, direct DNMT1 inhibitors without activity as DNA intercalators. Seven of these compounds exhibit modest selectivity for DNMT1 versus inhibition of DNMT3A/DNMT3L.

### Materials and Methods

#### DNMT Expression and Purification

Truncated forms of human DNMT1 (RFTS-lacking DNMT1, amino acids 621–1616 and RFTS-containing DNMT1, amino acids 351–1616) were expressed and purified as previously reported [27]. Full-length human DNMT3L was expressed and purified as previously described [30]. The catalytic domain of human DNMT3A (CD-DNMT3A; amino acids 611–912) was expressed as an N-terminally his-tagged protein in Rosetta 2(DE3)pLysS competent cells (Novagen). Cells were grown to an optical density of ~0.5 and then cooled to 18°C. Protein expression was induced with 0.5 mM IPTG and cultures were grown at 18°C for 16 hours. Following cell lysis, protein was purified via metal affinity using Ni-NTA resin (GE Life Sciences). Bound protein was eluted using a linear gradient from 0.2 to 1.25 M NaCl. Eluted CD-DNMT3A was concentrated and stored at −80°C in 50% glycerol. The CD-DNMT3A-DNMT3L complex was prepared by overnight 1:1 molar incubation at 4°C in 10 mM Tris pH 7.5, 300 mM NaCl, 2 mM DTT. All proteins were quantified using A280 and calculated extinction coefficients.

#### Endonuclease-Coupled DNA Methylation Assay

DNMT activity was measured at 37°C using a fluorogenic DNA methylation assay [27]. In short, a hemi-methylated hairpin oligonucleotide with a 5’ fluorophore and a 3’ quencher is methylated by addition of a DNMT and the methyl-donating co-factor, S-adenosyl methionine (SAM, HPLC-purified from Sigma). The fully methylated product is a substrate for the restriction endonuclease GlaI (Sibenzyme), which cleaves the oligonucleotide, releasing the fluorophore from the quencher and generating fluorescence. The oligonucleotide substrate (5’-FAM-CCTATGGGmCATCAGTTTTCTGATGmCGmCATAGG-3’-Iowa Black, in which mC denotes 5-methyldeoxycytidylate residues), termed 8006, was synthesized by Integrated DNA Technologies, Coralville, IA. 96 well plates were performed in Costar black half-area plates and read in a Biotek Synergy Neo plate reader. 384 well assays were performed in Nunc flat-bottomed black plates and read in a PerkinElmer EnVision plate reader. FAM fluorescence was measured using excitation and emission wavelengths of 485 nm and 520 nm, respectively.

#### High-Throughput Assay and Hit Validation

HTS against the Spectrum compound collection (Microsource, Gaylordsville, CT) was performed in 384 well plates. Using a Multiflo dispenser (Biotek), 75 µl of assay buffer (10.8 mM Tris-HCl pH 7.5, 1.08 mM MgCl2, 1.08 mM DTT, 108 mM potassium glutamate, 0.108 mg/ml BSA, 10.8 µM SAM, 21.6 mM oligonucleotide 8006, 5.4% glycerol) was dispensed into each well. 1 µl of each compound at 880 µM in DMSO (columns 3–22) or DMSO (columns 1, 2, 23 and 24) was added to assay plates using a Microlab Star liquid handling robot (Hamilton) and the plate was incubated at 37°C for 10 minutes. Following this warming step, 5 µl enzyme solution (either a DNMT1-containing solution of 20 mM Tris-HCl, pH 7.5, 300 mM NaCl, 2 mM DTT, 49 nM DNMT1 [amino acids 621–1616], 0.08 U/µl GlaI, 30% glycerol to columns 2–23 or a GlaI control solution of 20 mM Tris-HCl, pH 7.5, 300 mM NaCl, 2 mM DTT, 0.08 U/µl GlaI, 30% glycerol to columns 1 and 24) was added and the plate was again incubated at 37°C for 25 minutes. The final composition of the HTS assay was 11.2 mM Tris-HCl, pH 7.5, 1 mM MgCl2, 1.12 mM DTT, 100 mM potassium glutamate, 18.5 mM NaCl, 0.1 mg/ml BSA, 10 µM SAM, 20 nM oligonucleotide 8006, 6.85% glycerol, 0.4 µM GlaI, 3 nM DNMT1, 10.9 µM test compound, 1.23% DMSO. Each 384 well plate contained two columns of negative (n) and positive (p) controls for inhibitor activity. Columns 2 and 23 were the n controls with DMSO without test compound. Columns 1 and 24 were the p controls without DNMT1. Thus, 320 compounds were assayed per 384 well plate in columns 3–22.

Assay performance was assessed across the screen using the following parameters: the signal-to-noise (S/N) ratio = (m_S - µ_n)/SD_n, the signal-to-background (S/B) ratio = µ_p/µ_n, and the Z’-factor = 1− 3*S_D/SD_p, in which SD_D and SD_p are standard deviations, and µ_p and µ_n are means of the n and p control wells of each plate [31].

Data across the screen were normalized to the p (0% DNMT1 activity) and n (100% DNMT1 activity) controls on each plate. Compounds that resulted in at least a 5 SD reduction in observed DNMT1 activity (<41%) were considered potential hits and were re-examined. Validation assays (81 µl total volume, identical buffer conditions to those used in the HTS screen) were performed in triplicate in 96 well plates with FAM fluorescence measured over the course of 1 hour. A control containing GlaI in the absence of DNMT1 was subtracted from each assay condition. Corrected
assay traces were plotted and compared to DMSO-containing control traces in Prism (GraphPad Software, Inc).

**Differential Scanning Fluorimetry (DSF) Assay**

DSF [32] was used to assess the ability of the validated inhibitors to bind directly to DNMT1 in the absence of DNA and SAM and alter the observed melting temperature ($T_m$). Assays (25 µl) were conducted in triplicate in a Bio-Rad C1000 Thermal Cycler – CFX Real-Time System using the FRET channel and contained 50 mM HEPES pH 7.5, 150 mM NaCl, 2 µM DNMT1 (amino acids 621–1616), 5X Sypro Orange (Invitrogen Molecular Probes), 100 µM compound, and 1% DMSO. A DMSO control assay in the absence of compounds was also examined. Temperature was increased from 25 to 95°C by 0.5°C per minute. Fluorescence traces were exported and analyzed by fitting to the Boltzmann equation in Prism to determine the $T_m$.

**GlaI Counterscreen**

GlaI inhibitors were identified and excluded using a fluorescence-based assay. Duplicate assays (80 µl) were conducted in 96 well plates. Assays contained 10 mM Tris-HCl pH 7.5, 100 mM potassium glutamate, 1 mM MgCl$_2$, 1 mM DTT, 0.1 mg/mL BSA, 5% glycerol, 5 nM oligonucleotide 8007 (5'-FAM-CCTATGmCGmCATCAGTTTTCTGATGmCGmCATAGG-3''), 0.94% DMSO was used as a negative control and daunorubicin (Sigma) was used as a positive control. Percent activity was determined by comparing product formation to a DMSO containing control assay.

**Concentration-Dependence Experiments**

IC$_{50}$ values for each validated direct inhibitor were determined under identical assay conditions (10 µM SAM and 20 nM oligonucleotide 8006) using the endonuclease-coupled DNA methylation assay in triplicate (80 µl) in a 96 well format. Assays contained 0.8 U GlaI, 2 nM DNMT1 (amino acids 621–1616) and 0.94% DMSO with inhibitor concentration varied from 0 to 10 µM. FAM fluorescence was measured in a Biotek Neo plate reader over the course of 45 minutes. A control containing GlaI in the absence of DNMT1 was subtracted from each assay condition. Corrected assay traces were fitted in Prism and percent activity was determined by comparing to a DMSO containing control. IC$_{50}$ values were determined by fitting the percent activity data using a unity Hill slope in Prism.

**Compound Selectivity**

After identifying direct inhibitors of DNMT1 lacking the RFTS domain, inhibitor selectivity was assessed by examining inhibition of RFTS-containing DNMT1 (amino acids 351–1616), CD-DNMT3A/DNMT3L complex, and the bacterial cytosine methyltransferase from M. SsSI (New England Biolabs) using the endonuclease-coupled DNA methylation assay. Triplicate assays (100 µl) containing 0.25 mM SAM, 0.2 µM oligonucleotide 8006, 0.8 U GlaI, 20 mM methyletransferase, 20 µM inhibitor and 1% DMSO were conducted in 96 well plates. Following enzyme addition, assay plates were incubated at 37°C for 75 min and fluorescence was measured in a BioTek Neo plate reader. A control containing only GlaI was subtracted from each assay. SGI-1027, a non-selective DNMT inhibitor [25], was used as a positive control and 5-aza-2-cytidine (Sigma) was used as a negative control. Percent activity was determined by comparing product formation to a DMSO containing control assay.

**Results**

**High Throughput Screening**

Targeting epigenetic changes is a promising cancer therapy strategy as aberrant DNA methylation is closely related to initiation and progression of many cancers [12]. The link between DNMT1 hyperactivity and cancer is well established [12,33], making DNMT1 an important cancer drug target. For this reason, we set out to develop a pipeline to discover novel, direct small molecule inhibitors of DNMT1 activity. The sine qua non for small molecule screening is a robust assay. We previously refined an in vitro assay for DNMT1 that couples DNA methylation to fluorescence generation using the restriction endonuclease GlaI [27], allowing for an activity assay that is exquisitely sensitive. This assay utilizes a hemimethylated hairpin DNA substrate with a 5'-FAM fluorophore and a 3' quencher. The fully methylated product oligonucleotide is a substrate for the restriction endonuclease GlaI. Cleavage of the product DNA releases the fluorophore from the quencher and generates fluorescence. Using this endonuclease-coupled DNA methylation assay, we showed that an N-terminal deletion of sequences up to and including the RFTS domain, the first 620 amino acids of DNMT1, was sufficient to disrupt DNMT1 activity [27]. This de-repressed form of DNMT1, with a $k_{cat}/K_m$ of $10^6$ M$^{-1}$ s$^{-1}$ [27], has sufficient catalytic power to allow for facile identification of inhibitors using the fluorogenic assay.

To determine if the endonuclease-coupled DNA methylation assay is suitable for HTS, we first evaluated the effect of DMSO on the observed activity of the de-repressed form of DNMT1 (amino acids 621–1616). The presence of up to 5% DMSO, the highest value tested, has no effect on DNMT1 activity (Fig. S1), indicating that DMSO does not retard DNMT1 or inhibit the DNA methylation detection system. Next, to increase throughput, we wished to ensure the assay could be miniaturized to 384 well plates. Range-finding experiments at a variety of substrate concentrations, enzyme amounts, volumes and time were performed. Conditions were chosen in which the DNA substrate is at 20 nM, 10–20 times $K_{m, DNA}$ [27,34], to bias against selection.
of DNA-competitive inhibitors, SAM is at 10 μM, 5 times $K_{m,SAM}$ [34], to permit identification of SAM competitors, and reactions were scored 25 minutes post enzyme addition. Using DNMT1 plus GlaI endonuclease to represent 100% DNMT1 activity, and GlaI in the absence of DNMT1 to represent 0% DNMT1 activity, a Z’-factor of 0.66 was obtained for the miniaturized DNA methylation assay (Fig. 1). The S/B ratio (4.2) and S/N ratio (12.9) indicate that the assay is robust for HTS.

The miniaturized endonuclease-coupled DNA methylation assay was used to screen the Spectrum compound collection for inhibitors of DNMT1 activity. These 2320 compounds provide a wide-range of biological activities and structural diversity. The collection contains drug and drug-like synthetic compounds as well as natural products. The quality of the assay was assessed by determining the S/B ratio, S/N ratio and Z’-factor values of 4.6±0.4, 8.8±1.4 and 0.52±0.06 respectively (Fig. 2), screening of the Spectrum collection was successful. Compounds were picked as hits if resulting activities were greater than 5 standard deviations below the mean of the negative controls (Fig. 3). Of 2320 compounds examined, 57 (2.5%) were primary hits. Each of the 57 primary hits was re-tested in triplicate for DNMT1 inhibition using the endonuclease-coupled DNA methylation assay. For validation, time-dependent reactions traces for each candidate inhibitor and a DMSO control reaction were collected. 11 of 57 primary hits failed to inhibit DNMT1 (Table S1), whereas 46 compounds inhibited product formation by at least 40% (Table S1), giving an apparent hit rate of ~2%.

Identification of Direct DNMT1 Inhibitors

Direct DNMT1 inhibitors have the potential to be useful molecular probes and possibly leads for drug development. To ensure that the inhibitory effect exhibited by the validated HTS hits stems from a direct interaction with DNMT1, we employed DSF [32] to determine the observed melting temperature ($T_w$) of DNMT1 in the presence of each candidate inhibitor. Compounds that directly interact with target proteins in the absence of substrates frequently stabilize against thermal denaturation and shift the observed $T_w$ to right. Thus, comparing the $T_w$ observed in the presence of a small molecule to that observed in the presence of DMSO should allow for detection of direct inhibitors. Of 46 compounds tested, five could not be assayed by DSF. These compounds either strongly quenched the Sypro Orange fluorescence signal or were fluorescent and interfered with the assay. The majority of compounds assayed had no significant effect of the observed melting temperature (Table S2). Addition of 27 of the 41 successfully assayed compounds resulted in $T_w$ values within ~0.5°C of that observed for DNMT1 alone. However, addition of 12 compounds shifted the observed $T_w$ to the right by at least 0.9°C (Table S2), indicating that the compounds make a stabilizing interaction with the protein.

Though 12 compounds bind DNMT1 directly as indicated by DSF, we aimed to eliminate those compounds that interfere with an aspect of the endonuclease-coupled DNA methylation assay, making it difficult to determine their effect on DNMT1 activity. To assess the possibility that the small molecules inhibit GlaI or quench fluorescence of the 5’-FAM product, the 12 candidate DNMT1 inhibitors were counterscreened against GlaI in a fluorogenic assay using a fully CpG-methylated hairpin DNA substrate (Fig. S2). Addition of 10 of the 12 compounds resulted in activities ≥79% of that observed with DMSO alone (Table S3). However, addition of two compounds resulted in observed activities of <40%. Due to their interference with the coupling reaction used in the DNA methylation assay, these compounds were not considered DNMT1 inhibitors.

To further investigate inhibition of DNMT1 by the direct-binding compounds, a detergent test was performed. Inhibition of each compound was investigated in the presence and absence of 0.01% Triton X-100 to eliminate promiscuous molecules that inhibit by nonspecific aggregation of protein targets [35]. The presence of Triton X-100 did not reduce the inhibition observed for 9 compounds (Table S4). However, observed inhibition was almost completely lost in the presence of detergent for one candidate inhibitor. For this reason, this compound was excluded from further study.

Finally, while the compounds shift the observed $T_w$ of DNMT1 in the absence of DNA, indicating that they directly interact with the enzyme, we sought to ensure that the mechanisms of inhibition of these compounds are not due to DNA intercalation. To address this possibility, the compounds were added to a DNA-ethidium bromide mixture. If the compounds compete with ethidium bromide and intercalate into DNA, the fluorescence intensity of ethidium bromide will decrease. None of the validated direct hits were capable of decreasing ethidium bromide fluorescence, indicating that they are not strong DNA intercalators (Table S5). In comparison, addition of daunorubicin, a known DNA intercalator [36], significantly reduced fluorescence in this assay. Overall, the high throughput pipeline described resulted in the discovery of 9 direct inhibitors of DNMT1 enzymatic activity (Fig. 4) from 2320 compounds in the Spectrum collection.

Concentration-Dependence of Inhibition

To gauge the potency of the 9 resulting compounds, the concentration-dependence of inhibition was investigated using the endonuclease-coupled DNA methylation assay. Under identical conditions (10 μM SAM and 20 nM hairpin oligonucleotide 8006), each inhibitor was varied from 100 nM to 10 μM. The

![Figure 1. Z’-factor determination of HTS assay in 384 well plates.](image)
percent activity observed at each condition was determined by comparing to an uninhibited DMSO-containing control reaction. The 9 compounds examined exhibited IC50 values ranging from 300 nM to 11 μM (Fig. 5 and Table 1).

Inhibitor Selectivity

Most characterized DNMT inhibitors are not isozyme-selective. With three catalytically active DNMTs in humans and the finding that DNMT3A acts as a tumor suppressor [21], isozyme selectivity of DNMT inhibitors is important. The compounds discovered using this high throughput pipeline are direct inhibitors of the truncated and activated form of DNMT1 (amino acids 621–1616) in vitro. To investigate selectivity for inhibition of methylation of oligonucleotide 8006, three methyltransferases was examined: RFTS-containing DNMT1 (amino acids 351–1616), the CD-DNMT3A/DNMT3L complex, and the bacterial methyltransferase from M.SssI. SGI-1027 was used as a positive control because it has been shown to inhibit each enzyme [25]. 5-aza-cytidine was used as a negative control; this nucleoside inhibitor must be incorporated into DNA to exert inhibitory effects on DNMTs. The percent activity observed for each enzyme was determined by comparing to an uninhibited DMSO-containing control assay. As expected, SGI-1027 inhibited the activity of all DNMTs tested,
while addition of 5-aza-cytidine did not inhibit any enzyme examined (Table 2). Seven of nine compounds discovered in this HTS campaign showed some selectivity. Whereas triclosan (cmpd 13) and alizarin (cmpd 26) inhibited each enzyme to a similar degree, the other seven compounds either showed a preference for inhibition of DNMT1 or for inhibition of both DNMT1 and M.SssI, but not DNMT3A/DNMT3L. The observed selectivity for DNMT1 over DNMT3A/DNMT3L in this assay was two to four-fold.

---

**Figure 3. HTS screen of the Spectrum library.** Raw fluorescence data from the Spectrum screen were normalized to the assay controls on each plate. The normalized data are shown as circles with each assay plate being a different color. The solid lines represent the average value of the positive (▲) and negative (▼) controls. The dashed black lines represent one standard deviation from the positive and negative controls. The red dashed line represents 5 standard deviations below the average negative control. Compounds below this cutoff were considered primary hits. 57 of the 2320 compounds were identified as hits.

doi:10.1371/journal.pone.0078752.g003

**Figure 4. Structure of identified direct DNMT1 inhibitors.** Shown are the 9 identified inhibitors from the Spectrum collection. Compound numbers correspond to those in all tables and Figure 5. Five compounds contain a quinone-like substructure. Two identified inhibitors are chlorobenzene compounds.

doi:10.1371/journal.pone.0078752.g004
Drastic changes to normal DNA methylation patterns occur in malignancy. The genome typically becomes globally hypomethylated with regional hypermethylation and gene silencing of tumor suppressor genes [12]. Targeting these epigenetic changes is a promising cancer therapy strategy. While modified cytosine nucleosides, such as 5-aza-cytidine, have been approved to treat myelodysplastic syndrome, such demethylators are not DNMT isozyme specific and require incorporation into DNA to exert their effects [22], suggesting that there is room for improvement in probe development. Molecular probes targeted at DNMT1 enzyme activity will also be useful in furthering our understanding of cancer etiology. Such molecules can be used to determine the degree to which DNA methyltransferase enzyme activity as opposed to DNMT physical interactions is important for gene silencing and carcinogenesis. Direct inhibitors may also serve as lead compounds for new cancer therapeutics. Thus, the objective of developing this high throughput pipeline was to identify novel direct inhibitors of DNMT1 activity.

The HTS assay described herein is ultrasensitive and requires only 0.24 pmol of DNMT1 (amino acids 621–1616) per well. In comparison, a recently described scintillation proximity assay [29] required almost 0.9 pmol of a similar DNMT1 construct. In contrast, a different endonuclease-coupled assay that uses biotin to attach a DNA substrate to a microtiter plate [37] used nearly 17 pmol of full-length DNMT1 per reaction. Our fluorescence-based DNA methylation assay for DNMT inhibitors requires little manipulation. The assay can be accomplished by simply adding test compounds and enzyme to assay solution and reading the fluorescence generated after a short 37°C incubation. Less than 1 hour is required from plate set-up to final reading, making the assay simple and scalable.

The initial screen of the Spectrum collection yielded 57 hits that inhibited DNMT1 activity by at least 5 standard deviations below the negative control. Following hit validation, 46 compounds reliably inhibited fluorescence-generation in the endonuclease-coupled DNA methylation assay. The validated inhibitors were examined for their ability to bind directly to DNMT1 using DSF [32]. These assays can be performed in 96 well and 384 well plates in a real-time PCR machine, permitting HTS for direct inhibition. Twelve compounds were determined to bind directly with DNMT1 on account of their ability to shift the observed $T_{m}$ of at least 0.9°C. A shift of the observed $T_{m}$ of at least 0.9°C was used as cutoff for direct binders. This is in line with previous studies that have shown changes in $T_{m}$ of 1–2°C or greater for direct binders of other

**Table 1.** IC$_{50}$ values of the 9 compounds identified using the high throughput pipeline.

| Cmpd # | Assay Plate | Well ID | Cmpd ID | IC$_{50}$ (µM)* |
|--------|-------------|---------|---------|-----------------|
| 13     | 3           | F8      | 01505465 | 3.8 (2.4–5.9)   |
| 22     | 4           | G5      | 01504078 | 0.47 (0.33–0.67)|
| 24     | 4           | J5      | 01503867 | 0.40 (0.27–0.60)|
| 26     | 5           | B22     | 00210850 | 11 (7.9–15)     |
| 29     | 5           | M9      | 01505786 | 6.1 (3.4–11)    |
| 33     | 6           | A4      | 01505143 | 2.9 (1.7–5.0)   |
| 36     | 6           | G15     | 01504080 | 0.45 (0.33–0.60)|
| 40     | 6           | K10     | 01505847 | 2.1 (1.7–2.6)   |
| 51     | 7           | O19     | 00201507 | 0.33 (0.22–0.51)|

*IC$_{50}$ values obtained by fitting percent activity data shown in Fig. 5 (with 95% confidence interval).
targets [32, 38, 39]. Of the 12 compounds that stabilize DNMT1 against thermal denaturation, 10 were further confirmed as lacking in GlaI inhibition and 9 were confirmed for their ability to inhibit in the presence of detergent. These secondary and confirmatory assays reduced the number of direct DNMT1 inhibitors to nine, 0.39% of all compounds screened.

All 9 inhibitors discovered using this high throughput pipeline show concentration-dependent inhibition of DNMT1 activity in vitro, with IC_{50} values ranging from 0.3–11 μM (Fig. 5 and Table 1). The majority of compounds identified are polycyclic aromatics (Fig. 4). Though many such compounds are DNA intercalators, these molecules did not inhibit Gla cleavage of the DNMT1 product oligonucleotide, they shifted the melting temperature of DNMT1 in DSF assays that included no DNA, and they failed to compete with ethidium bromide in DNA intercalation assays. Compounds similar in structure to our identified compounds, such as nitroflavones and dichlone [28], were recently reported to inhibit DNMT3A, suggesting that polycyclic aromatics may profitably be screened for inhibition of DNMT isoymes.

Our high throughput pipeline used a truncated form of DNMT1 (amino acids 621–1616) due to its increased in vitro activity [27]. To examine the ability of the identified compounds to inhibit full-length DNMT1, we tested them against RFTS-containing DNMT1 (amino acids 351–1616), a protein that behaves similarly to the full-length enzyme in vitro [27]. All of the identified compounds inhibited the activity of RFTS-containing DNMT1 (Table 2). To examine isoyme selectivity, the compounds were also screened against the CD-DNMT3A/DNMT3L complex and the bacterial DNMT from M.Sssl. Interestingly, only alizarin (cmpd 26) and triclosan (cmpd 13) appear to be nonselective. The seven other compounds identified in the HTS campaign exhibited two to four-fold selectivity for DNMT1 over DNMT3A/DNMT3L. While these initial finding are promising, further work is needed to characterize the new inhibitors. Structure activity relationship analyses of the inhibitors and related compounds could reveal more potent and specific DNMT1 inhibitors.

Of the compounds identified in our screen, five contain anthracene or anthraquinone-related structures. Laccaic acid A (LCA, cmpd 40), a highly substituted anthraquinone natural product, is ~5-fold more potent than alizarin (cmpd 26), an anthraquinone with only two hydroxyl substituents, suggesting that substituted anthraquinones represent a novel pharmacophore for DNMT1 inhibitors. LCA exhibited ~4-fold selectivity for DNMT1, while alizarin inhibited all DNMTs examined equally. Further characterization of LCA has shown that it is a DNA-competitive inhibitor, which reactivates expression of a set of methylation-silenced genes in MCF-7 breast cancer cells [34]. In addition, LCA reverses DNMT1-dependent oncogenic transformation and apoptosis in murine Rgs6−/− mouse embryonic fibroblasts [40].

This screen also yielded two chlorobenzene compounds, triclosan (cmpd 13) and triclabendazole (cmpd 29), with similar potencies against the activated form of DNMT1. A recent study has shown that treatment with triclosan reduced the levels of DNA methylation in HepG2 cells [41]. Triclabendazole but not triclosan exhibited selectivity for DNMT1 versus DNMT3A/DNMT3L.

The high throughput pipeline described in this study was used successfully to identify direct inhibitors of DNMT activity in vitro from a small chemical library. Whereas alizarin and triclosan appear to be nonspecific DNMT inhibitors, seven other compounds appear to be at least partially selective for DNMT1 over DNMT3A/DNMT3L. Ongoing experiments are designed to determine their mechanisms of inhibition, cellular availability and cellular isoyme specificity. Structure activity relationship data and co-crystallization studies are expected to aid in further defining DNMT1 pharmacophores. The pipeline described herein can be used to screen larger and more diverse libraries of chemical matter to discover additional tool compounds and leads for clinical DNMT1 inhibition.

### Supporting Information

**Figure S1** DMSO tolerance of DNMT1. DNMT1 activity was assayed at 20 nM oligonucleotide 8006 and 10 μM SAM using 2 nM DNMT1. DMSO concentration was varied from 0–5% (0– filled circles; 0.5– filled squares; 1– filled triangles; 2– filled diamonds; 3– open circles; 4– open squares; 5– open triangles). Addition of DMSO has little effect on the observed activity of DNMT1. RFU, relative fluorescence unit. (TIF)

**Figure S2** GlaI counterscreen. The effect of each compound on GlaI activity, the coupling enzyme used in the DNA methylation assay, was investigated using an internally quenched

| Compound # | RFTS-containing Dnmt1 | DNMT3A/DNMT3L Complex | M.Sssl DNMT |
|------------|------------------------|------------------------|-------------|
| 13         | 25 ± 8                 | 23 ± 10                | 25 ± 4      |
| 22         | 40 ± 10                | 88 ± 11                | 20 ± 5      |
| 24         | N.D.*                  | 49 ± 16                | 49 ± 12     |
| 26         | 51 ± 10                | 49 ± 12                | 68 ± 11     |
| 29         | N.D.                   | 110 ± 5                | 19 ± 6      |
| 33         | 38 ± 13                | 74 ± 13                | 86 ± 9      |
| 36         | 30 ± 12                | 83 ± 15                | 40 ± 7      |
| 40         | 21 ± 9                 | 80 ± 10                | 82 ± 10     |
| 51         | 24 ± 11                | 61 ± 12                | 36 ± 7      |
| SGI-1027   | N.D.                   | N.D.                   | N.D.        |
| 5-aza-cytidine | 100 ± 11               | 97 ± 10                | 100 ± 5     |

*N.D. – No detectable activity observed above background.

doi:10.1371/journal.pone.0078752.t002
DNMT1-Targeted HTS Pipeline

hairpin DNA with a fully methylated GGGC site (the cleavage site of GluL). GluL cleavage of the oligonucleotide releases the 5′ fluorophore from the 3′ quencher, generating fluorescence in real-time. Shown is the time-dependent cleavage of 5 nM oligonucleotide substrate 9007 with 0.2 U/l of enzyme in the presence of DMSO (black), or 11 µM of each compound (13– red; 22– blue; 24– green; 26– purple; 29– red; 30– blue; 33– green; 36– purple; 40– red; 44– blue; 51– green; 53– purple).

Table S1 Validation of the initial 57 hits from the Spectrum HTS assay. Initial hits were validated as DNMT1 inhibitors using the endonuclease-coupled DNA methylation assay. Each compound was assayed in triplicate. Shown is the observed melting temperature (Tm) of DNMT1 in the presence and absence of validated hits. 12 compounds stabilized DNMT1 against thermal denaturation and shifted the observed Tm to right by at least 0.9°C, indicating that they bind directly to DNMT1.

Table S2 Melting temperature of DNMT1 determined using DSF. DSF was used to determine the observed melting temperature (Tm) of DNMT1 in the presence and absence of validated hits. 12 compounds stabilized DNMT1 against thermal denaturation and shifted the observed Tm to right by at least 0.9°C, indicating that they bind directly to DNMT1.

Table S3 Effect of compounds on GluL endonuclease activity. A GluL counterscreen was performed to determine if the compounds inhibit the restriction enzyme used in the DNA methylation assay. Two of the twelve compounds that shifted the melting temperature of DNMT1 inhibited GluL activity in this assay. These compounds were not studied further.

Table S4 Effect of detergent of inhibition. The percent activity observed using 5 µM compound in the presence and absence of 0.01% Triton X-100 was determined. The observed inhibition with compound 44 was sensitive to detergent. The inhibitory effect of the other nine compounds examined was not sensitive to detergent.

Table S5 DNA Intercalation Assay. DNA intercalation activities of candidate inhibitors were assessed using an assay containing calf thymus DNA and ethidium bromide. Ethidium bromide fluorescence was measured using excitation and emission wavelengths of 320 and 600 nm, respectively. Compounds that intercalate DNA decrease the observed fluorescence. Daunorubicin, a known DNA intercalator, was used as a positive control and significantly reduced the fluorescence signal. None of the compounds identified in the HTS campaign had a significant effect on observed fluorescence, indicating that they do not intercalate into DNA under reaction conditions.

Author Contributions
Conceived and designed the experiments: RLF MW CB. Performed the experiments: RLF. Analyzed the data: RLF MW CB. Contributed reagents/materials/analysis tools: MW FC. Wrote the paper: RLF CB.

References

1. Jurkowski RZ, Jurkowski TP, Jelsch A (2011) Structure and function of mammalian DNA methyltransferases. ChemBiochem 12: 206-222.

2. Goyal R, Reinhardt R, Jelsch A (2006) Accuracy of DNA methylation pattern preservation by the Dnm1 methyltransferase. Nucleic Acids Res 34: 1182-1188.

3. Yoder JA, Soman NS, Verdine GL, Bestor TH (1997) DNA (cytosine-5)-methyltransferases in mouse cells and tissues. Studies with a mechanism-based probe. J Mol Biol 270: 385-395.

4. Bird A (2002) DNA methylation patterns and epigenetic memory. Genes Dev 16: 6-21.

5. Gavrilidi T, Sharma RP (2010) Histone modifications, DNA methylation, and schizophrenia. Neurosci Biobehav Rev 34: 682-688.

6. Tsankova N, Renthal W, Kumar A, Nestler EJ (2007) Epigenetic regulation in schizophrenia. Nat Rev Neurosci 8: 355-367.

7. Karouzakis E, Gay RE, Gay S, Neidhart M (2009) Epigenetic control in rheumatoid arthritis synovial fibroblasts. Nat Rev Rheumatol 5: 266-272.

8. Mishra N, Brown DR, Olorenshaw IM, Kammer GM (2001) Trichostatin A reverses skew expression of CD154, interleukin-10, and interferon-gamma in psychiatric disorders. Nat Rev Neurosci 2: 1-8.

9. Klein CJ, Botuyan MV, Wu Y, Ward CJ, Nicholson GA, et al. (2011) Mutations in DNMT1 cause hereditary sensory neuropathy with dementia and hearing loss. Nat Genet 43: 595-600.

10. Szyf M (1997) DNA methylation patterns and their role in mammalian development. Annu Rev Genet 31: 227-261.

11. Tabery M, Jones PA (2011) DNA methylation and cancer. Prog Drug Res 67: 197-252.

12. Cavenee WK, Gajewski TF, Sidransky D, et al. (2001) DNA methylation: a mechanism for cellular transformation. Annu Rev Pathol 6: 37-75.

13. Pardini MN, Thiele CJ, Jentsch D, et al. (2011) Interplay of cell cycle and chromatin remodeling in cellular transformation. Nat Rev Mol Cell Biol 12: 513-525.

14. MacLeod AR, Boulet A, Szyf M (1995) Regulation of DNA methylation by the Ras signaling pathway. J Biol Chem 270: 11327-11337.

15. Slack A, Cervoni N, Pinard M, Szyf M (1999) DNA methyltransferase is a downstream effector of cellular transformation triggered by simian virus 40 large T antigen. J Biol Chem 274: 10105-10112.

16. Doherty C, Vogel AE, Laidi PW (2000) Complete genetic suppression of polyp formation and reduction of CagA and hypermethylation in Apo(Min/+)+ Dnm1-hypomorphic Mice. Cancer Res 62: 1296-1299.

17. MacLeod AR, Szyf M (1995) Expression of antisense to DNA methyltransferase mRNAs induces DNA demethylation and inhibits tumorigenesis. J Biol Chem 270: 8037-8043.

18. Ramachandran S, MacLeod AR, Pinard M, von Hoef E, Szyf M (1997) Inhibition of tumorigenesis by a cytosine-DNA methyltransferase, antisense oligodeoxynucleotide. Proc Natl Acad Sci U S A 94: 684-689.

19. Ley TJ, Ding L, Walter MJ, McLellan MD, Langerod C, et al. (2010) DNMT1 mutations in acute myeloid leukemia. N Engl J Med 362: 2424-2433.

20. Koo BC, Jones PA (2006) Epigenetic therapy of cancer: past, present and future. Nat Rev Drug Discov 5: 37-50.

21. Stromberg C, Lyko F (2008) Modes of action of the DNA methyltransferase inhibitors azacytidine and decitabine. Int J Cancer 129: 8-13.

22. Gros C, Fasy J, Halley L, Dulau I, Erdmann A, et al. (2012) DNA methylation inhibitors in cancer: recent and future approaches. Biochimie 94: 2280-2296.

23. Dutta J, Ghoshal K, Denny WA, Gamage SA, Brodie DG, et al. (2009) A new class of quinoline-based DNA hypomethylating agents reactivates tumor suppressor genes by blocking DNA methyltransferase 1 activity and inducing its degradation. Cancer Res 69: 4277-4285.

24. Eldem RM, Reisine T (2011) Screening for compounds that modulate epigenetic regulation of the transcriptome: an overview. J Biomed Screen 16: 1137-1152.

25. Sveda F, Fagan RL, Wean M, Avvakumov GV, Walker JR, et al. (2011) The replication focus targeting sequence (RFTS) domain is a DNA-competitive inhibitor of Dnm1. J Biol Chem 286: 13544-13551.

26. Ceccaldi A, Rajaveedu A, Ragout S, Sennaaz-Braunfort C, Bashkryev P, et al. (2013) Identification of novel inhibitors of DNA methylation by screening of a chemical library. ACS Chem Biol 8: 543-548.

27. Kalgrove JA, Du X, Melito L, Wei S, Wang C, et al. (2011) Identification of DNMT1 selective antagonists using a novel scintillation proximity assay. J Biol Chem 286: 19673-19680.
30. Kareta MS, Botello ZM, Ennis JJ, Chou C, Chedin F (2006) Reconstitution and mechanism of the stimulation of de novo methylation by human DNMT3L. J Biol Chem 281: 25893–25902.

31. Zhang JH, Chung TD, Oldenburg KR (1999) A Simple Statistical Parameter for Use in Evaluation and Validation of High Throughput Screening Assays. J Biomol Screen 4: 67–73.

32. Niesen FH, Berglund H, Vedadi M (2007) The use of differential scanning fluorimetry to detect ligand interactions that promote protein stability. Nat Protoc 2: 2212–2221.

33. Teodoridis JM, Hardie C, Brown R (2008) CpG island methylator phenotype (CIMP) in cancer: causes and implications. Cancer Lett 268: 177–186.

34. Fagan RL, Cryderman DE, Kopelovich L, Wallrath LL, Brenner C (2013) Laccaic Acid A Is a Direct, DNA-competitive Inhibitor of DNA Methyltransferase 1. J Biol Chem 288: 23858–23867.

35. Feng BY, Shoichet BK (2006) A detergent-based assay for the detection of promiscuous inhibitors. Nat Protoc 1: 550–553.

36. Hortobagyi GN (1997) Anthracyclines in the treatment of cancer. An overview. Drugs 54 Suppl 4: 1–7.

37. Ceccaldi A, Rajavelu A, Champion C, Rampon C, Jurkowska R, et al. (2011) C3-DNA methyltransferase inhibitors: from screening to effects on zebrafish embryo development. ChemBioChem 12: 1337–1345.

38. Wan KF, Wang S, Brown CJ, Yu VC, Entzeroth M, et al. (2009) Differential scanning fluorimetry as secondary screening platform for small molecule inhibitors of Bel-4X. Cell Cycle 8: 3943–3952.

39. Amaning K, Lewinski M, Valler F, Steier V, Marciroca C, et al. (2013) The use of virtual screening and differential scanning fluorimetry for the rapid identification of fragments active against MEK1. Bioorg Med Chem Lett 23: 3620–3625.

40. Huang J, Stewart A, Maity B, Hagen J, Fagan RL, et al. (2013) RGS6 suppresses Ras-induced cellular transformation by facilitating Tip60-mediated Dnmt1 degradation and promoting apoptosis. Oncogene, In Press.

41. Ma H, Zheng L, Li Y, Pan S, Hu J, et al. (2013) Tricosanol reduces the levels of global DNA methylation in HepG2 cells. Chemosphere 90: 1023–1029.