A High Throughput Assay for Discovery of Bacterial β-Glucuronidase Inhibitors

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Abstract: CPT-11 is a widely-used anti-cancer drug that is converted in vivo to its active metabolite, SN-38. In the liver, enzymes detoxify SN-38 by coupling it to a glucuronidate moiety and this inactive compound (SN-38G) is excreted into the gastrointestinal tract. In the intestine, commensal bacteria convert the SN-38G back to the active and toxic SN-38 using bacterial β-glucuronidase (GUS). This intestinal SN-38 causes debilitating diarrhea that prevents dose-intensification and efficacy in a significant fraction of patients undergoing CPT-11 treatment for cancer. This CPT-11 metabolic pathway suggests that small molecule inhibitors of GUS may have utility as novel therapeutics for prevention of dose-limiting diarrhea resulting from CPT-11 therapy. To identify chemical inhibitors of GUS activity, we employed and validated a high throughput, fluorescence-based biochemical assay and used this assay to screen a compound library. Novel inhibitors of GUS were identified with IC50 values ranging from 50 nM to 4.8 μM. These compounds may be useful as chemical probes for use in proof-of-concept experiments designed to determine the efficacy of GUS inhibitors in altering the intestinal metabolism of drugs. Our results demonstrate that this high throughput assay can be used to identify small molecule inhibitors of GUS.

Keywords: β-glucuronidase, CPT-11, screen, inhibitor.

INTRODUCTION

Camptothecin is a potent anti-cancer compound that was originally purified from the tree Camptotheca acuminata and whose structure was elucidated in 1966 [1]. It inhibits the catalytic cycle of human topoisomerase I, which regulates the superhelical tension associated with DNA replication and is preferentially active in rapidly proliferating cells [2, 3]. In preliminary human studies, camptothecin exhibited significant toxicity and poor bioavailability [4]. The camptothecin derivatives topotecan and CPT-11 (also called irinotecan) have since been discovered to have improved toxicity and bioavailability properties and these drugs are now in clinical use as anti-cancer therapies [4], although they still generate debilitating side effects in cancer patients that limit efficacy. CPT-11 is one of the three commonly-used chemotherapeutic agents for colon, lung, and brain cancer and it has also been used against refractory forms of leukemia and lymphoma [5]. CPT-11 is a prodrug, having a carbamate-linked dipiperidino group that significantly increases its solubility and bioavailability [4]. This dipiperidino group is removed in vivo by carboxylesterases to produce the active metabolite, SN-38 [6].

The dose-limiting side effect of CPT-11 is severe diarrhea generated by its complex activation and metabolism [7, 8]. SN-38, the active metabolite of CPT-11, is glucuronidated in the liver by UDP-glucuronosyltransferase (UGT) enzymes [9] resulting in the inactive SN-38G [10], which is excreted via the biliary ducts into the gastrointestinal (GI) tract. Once in the intestines, SN-38G serves as a substrate for bacterial β-glucuronidase (GUS) enzymes that are produced by bacteria normally inhabiting the intestines. The removal of the glucuronide group by GUS generates a carbon source for the bacteria and, in the process SN-38G is reactivated back to the active and toxic SN-38 [11, 12]. This reactivated SN-38, now present in the intestinal lumen, is toxic to the intestinal cells resulting in delayed diarrhea that prevents dose-intensification and efficacy in a significant fraction of patients undergoing CPT-11 treatment for cancer [13, 14].

The concept of using antibiotics to reduce GI bacteria levels to prevent the regeneration of SN-38 in the intestine has been examined [15]. However, the use of antibiotics has several disadvantages. Intestinal bacteria function in essential pathways in carbohydrate metabolism, vitamin production, and the processing of bile acids, sterols and xenobiotics [16, 17]. Thus, the removal of GI bacteria is not recommended for patients already challenged by cancer and chemotherapy. In addition, elimination of symbiotic GI flora increases the chances of infections by pathogenic bacteria, including enterohemorrhagic E. coli and Clostridium difficile [18-24].

β-glucuronidase enzymes hydrolyze glucuronic acid sugar moieties from a variety of compounds [25]. The pres-
ence of GUS in a wide range of bacteria is exploited to detect bacterial contamination in commonly-used water purity tests [26]. Prior to our work, only relatively weak inhibitors of β-glucuronidases with \( K_i \) values ranging from 25 μM to 2 mM have been described [27, 28]. Thus, we have sought to identify potent and selective inhibitors of bacterial β-glucuronidases to inhibit the generation of SN-38 in the intestines and thus reduce or eliminate the GI toxicity of CPT-11 treatment without killing the useful bacteria required for intestinal health. From the high throughput screen for GUS inhibitors that is described in this report, we selected four hits for follow-up studies. We have recently published the results of these studies [29]. These compounds showed complete selectivity for bacterial GUS versus the homologous mammalian enzyme. The compounds demonstrated GUS inhibitory activity in living bacteria with IC50 values ranging from 18 nM to 1.3 μM with no effect on bacterial viability even at 100 μM. Furthermore, oral administration of one of these inhibitors protected mice from CPT-11-induced diarrhea.

In this report, we describe the development and validation of the high throughput GUS activity assay that led to the discovery of small molecule inhibitors of GUS activity. An existing GUS activity assay using the fluorogenic substrate 4-methylumbelliferyl glucuronide (4MUG) was modified and formatted for HTS. In this assay, the non-fluorescent 4MUG is hydrolyzed by GUS generating 4-methylumbelliferon, a highly fluorescent molecule. This optimized and validated assay was then used to screen a chemical library in search of inhibitors of GUS.

**MATERIALS AND METHODS**

**Reagents**

All common reagents such as HEPES, Triton X-100, D-Glucaric acid-1,4-lactone and dimethyl sulfoxide (DMSO) were reagent-grade quality obtained from Thermo Fisher Scientific (Waltham, MA) or Sigma-Aldrich (St. Louis, MO). 4-methylumbelliferyl glucuronide (4MUG) was obtained from Sigma. The solid black 384-well plates (cat# M) with no effect on bacterial viability even at 100 μM. Furthermore, oral administration of one of these inhibitors protected mice from CPT-11-induced diarrhea.

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**GUS Assay**

The GUS assay was performed by the addition of 0.5 μl of compound (or DMSO) to the well of a black 384-well plate followed by the addition of 30 μl of diluted GUS enzyme (83 pm GUS). The enzyme was diluted into 50 mM HEPES, pH 7.4 and 0.017% Triton X-100. After addition of enzyme, the reaction was initiated by addition of 20 μl of 4MUG substrate (312.5 μM 4MUG) diluted into 50 mM HEPES, pH 7.4. 4MUG stock solutions were made in the same buffer. Final buffer in the assembled assay was 50 mM HEPES, pH 7.4 and 0.01% Triton X-100. Following the initiation of the reaction with 4MUG, the plates were incubated for 30 minutes at 23°C followed by the addition of 20 μl of 1 M sodium carbonate solution to stop the reaction. The GUS, 4MUG, and stop solutions were delivered to the plate using Multidrop™ liquid handlers (Thermo). The plate was then read in the Victor V plate reader. The fluorescence was measured using the 355 nm excitation filter and 460 nm emission filter (the umbelliferone filter set). Read time was 0.1 s/well and fluorescence expressed in relative fluorescence units (RFU). Error bars for data points represent standard deviations. Z'-factor values were derived by the published formula with Z'-factors of 0.5 – 1.0 indicating that an assay is robust for screening [30].

**High Throughput Screen**

For the high throughput screening of the Asinex chemical library, 0.5 μl of 1 mM compound in 100% DMSO was added to empty 384-well plates using a Biomek NX (Beckman Coulter Inc., Fullerton, CA), resulting in a final compound concentration of 10 μM for the primary screen. This dry spotting of compounds into assay plates was performed as a routine method of conserving compound stock solutions. This spotting method has been validated for transfer of 0.5 μl DMSO with a CV of 6.3% and 98% accuracy. Final assay concentrations in the 50 μl enzyme reaction were 50 pM GUS, 125 μM 4MUG, 10 μM compound, 1% DMSO (from compound). For the screening, each plate had maximum (max) and minimum (min) controls with DMSO spotted in the wells. Max signal positive control wells contained only DMSO (no compound), while the min signal control wells were obtained by adding enzyme dilution buffer instead of GUS. D-Glucaric acid-1,4-lactone would not dissolve in DMSO or water at a high enough concentration to generate the min controls. The percent inhibition was calculated based on the max and min controls on the plate. The definition of a hit was a compound that displayed ≥50% inhibition in the primary screen. IC50 value was defined as the concentration of inhibitor calculated to inhibit 50% of the assay signal based on a serial dilution of compound. Values were calculated using either a four or three-parameter dose response (variable slope) equation in Graphpad Prism or ActivityBase (IDBS, Alameda, CA). For the IC50 determinations, serial dilutions of compounds were performed in 100% DMSO with a two-fold dilution scheme resulting in 10 concentrations of compound starting at a high concentration of 20 μM compound in the final assay. For the HTS assay, these compound dilutions were spotted onto plates and the assay performed at 1% DMSO final concentration as described for the primary screen.

**RESULTS**

Bacterial GUS activity can readily be measured using the substrate 4-methylumbelliferyl glucuronide (4MUG) [26]. Upon hydrolysis of 4MUG by GUS, the fluorescent compound 4-methylumbelliferon (4MU) is generated. We used this substrate to detect GUS activity in an end point assay that is amenable for high throughput screening for inhibitors of GUS. The assay was developed with a final volume of 50 μl using the 384-well plate format to match the intended screening format. The general assay set-up included the addition of diluted GUS enzyme to the well followed by the addition of 4MUG to initiate the reaction (Fig. 1). The reaction was terminated with the addition of a 1 M Na2CO3 solution.
and the fluorescence measured in a plate reader. The final assay buffer consisted of 50 mM HEPES pH 7.4 and 0.01% TX-100. The $K_m$ for 4MUG was experimentally determined to be 125 μM (data not shown) and therefore this was the concentration of 4MUG used in this assay. Stopping of the reaction was confirmed by time course studies following termination of the assay (data not shown).

**Fig. (1).** Schematic of the GUS high throughput enzyme assay.

The activity in the assay was directly proportional to the concentration of GUS, up to at least 100 pM enzyme (Fig. 2). We chose to use 50 pM as the final concentration in the assay to provide a robust assay window. This concentration of enzyme resulted in a linear time course in this assay for 40 minutes under final assay conditions in the presence of 1% DMSO (Fig. 3). The enzyme titration and time course data suggested that the assay conditions should be sensitive to compounds that are competitive inhibitors of GUS activity.

**Fig. (2).** GUS enzyme titration. GUS was titrated into the assay under final assay conditions with an incubation time of 30 min in the presence of 1% DMSO. Data points represent the average of three determinations per concentration and error bars represent standard deviations. Data are representative of three independent experiments.

Since compounds to be screened are dissolved in DMSO, the tolerance for DMSO was examined (Fig. 4). The assay was shown to be tolerant to DMSO concentrations up to 4%. In the final screening assay, the maximal concentration that the enzyme is exposed to is 2% DMSO, before the addition of substrate, after which the final DMSO was set at 1%. All experiments in this report were performed in the presence of 1% DMSO.

**Fig. (3).** GUS assay time course. A time course study was performed under final assay conditions in the presence of 1% DMSO. Data points represent the average of three determinations per time point and error bars represent standard deviations. Data are representative of three independent experiments.

The variability of the assay was characterized in a 384-well plate format. The HTS version of the assay employed a work-station level automated liquid handling system using the Biomek NX for delivery of compound or DMSO (0.5 μl) to the plate followed by addition of GUS (30 μl), 4MUG (20 μl) and Na$_2$CO$_3$ solution (20 μl), in that order, using the Multidrop™ liquid handler (Thermo). To assess single-point variability of the assay with full automation, all wells of a 384-well plate were pre-spotted with 0.5 μl of DMSO (Fig. 5). The pre-spotted DMSO plates simulated compound plates.
that were to be screened. Two plates for each condition were used to determine the maximum (max) signal and minimum (min) signal. The %CV of the max and min plates were 2.6 and 17.5%, respectively. The Z’-factor for the inhibition assay window was 0.81. D-Glucaric acid-1,4-lactone has been reported to inhibit GUS [31]. The IC₅₀ value for this control inhibitor was determined using the HTS protocol, including use of automation (Fig. 6). The IC₅₀ values obtained were 21 and 17 μM for two separate determinations. Solubility of this control inhibitor limited the use of concentrations >100 μM, hence partial IC₅₀ curves were obtained. Despite this limitation, these values are in good agreement with the published Ki value of 25 μM [31]. The Hill slopes were both 1.07, very close to the expected value of 1.0 for inhibition of a single enzyme. Thus, reproducible IC₅₀ values can be obtained using the automated version of the GUS assay. This data indicated that the assay was highly robust with minimal variability and amenable for high throughput screening for inhibitors of GUS activity.

Fig. (5). GUS assay variability assessment. All wells of a 384-well plate were pre-spotted with 0.5 μl of DMSO. Two plates each were used to determine the maximum signal (■) and minimum signal (●).

Fig. (6). IC₅₀ value determinations using the automated GUS assay. Data points represent the average of three determinations per concentration and error bars represent standard deviations. Data are representative of two independent experiments.

As part of assay validation, a small collection of 1,120 compounds purchased from the Prestwick Chemical company was screened to assess the performance of the assay in the presence of diverse compounds using the HTS protocol (Fig. 7). The Prestwick collection of compounds was screened at 10 μM compound concentration. For this assay, the inhibitor cut-off was defined as 50% inhibition based on plate controls. Due to solubility limitation of inhibitor, we obtained the min controls on screening plates by leaving the enzyme out of these wells (buffer alone was added). The fluorescence obtained by leaving out the enzyme was the same as that of completely inhibited enzyme (data not shown). We obtained 40 actives (actives were defined as those compounds demonstrating ≥50% inhibition) from this test set screening resulting in a hit rate of 3.6%. The Z’-factors of the controls for each of the four plates were all ≥0.8.

Fig. (7). Screen of the Prestwick collection with the GUS assay. Each point represents a compound. The percent inhibition values were calculated relative to controls on the plates.

A diverse collection of 11,520 small organic molecules were purchased from Asinex Corporation and screened at 10 μM compound concentration for activity in this GUS assay. The plate Z’-factors for the screen were all ≥0.82. A total of 583 actives (≥50% inhibition) were obtained for an initial active rate of 5.1%. We sought to limit follow-up confirmatory IC₅₀ determinations to 120 actives, so the active cut-off was raised to ≥89% inhibition to obtain the most potent actives. Of these 120 compounds tested in IC₅₀ assays, 93 of them (77.5%) confirmed with an IC₅₀ of <20 μM. The Hill slopes generated from such data can be used to triage hits since steep slopes, i.e. significantly greater than 1.0, can be an indication of compound aggregation which is a non-specific inhibition mechanism displayed by some compounds [32-33]. Of the 93 confirmed hits, 26 had Hill slopes between 0.8 and 1.2. The potencies of these hits ranged from 50 nM to 4.8 μM. The Hill slopes from Table 1 indicated that 3 of these (compounds 1, 4 and 5) are structurally related to the quinolinone thiourea scaffold previously published [29]. Here we show the structures of 8 more novel GUS inhibitors as representative examples of the hits obtained from the screen with IC₅₀ values ranging from 1.7 to 4.8 μM (Table 1). The concentration response data used to calculate IC₅₀ values for two compounds are provided as representative curves (Fig. 8). Examination of the structures from Table 1 indicated that 3 of these (compounds 1, 4 and 5) are structurally related to the quinolinone thiourea scaffold previously published [29]. However, compounds 2, 3 and 6 - 8 are novel and structurally unique inhibitors of GUS.
| Compound | Asinex Identifier * | Structure | $IC_{50}$ (μM) $^b$ |
|----------|---------------------|-----------|------------------|
| 1        | ASN 03367547        | ![Structure](structure1.png) | 1.7              |
| 2        | ASN 03795365        | ![Structure](structure2.png) | 1.9              |
| 3        | BAS 06980438        | ![Structure](structure3.png) | 1.9              |
| 4        | ASN 03272623        | ![Structure](structure4.png) | 2.8              |
| 5        | ASN 03776465        | ![Structure](structure5.png) | 3.0              |
| 6        | BAS 00288912        | ![Structure](structure6.png) | 3.2              |

* identifiers correspond to ChemDiv catalog number **a**

$^b$ Values are IC50 (μM), indicating the concentration at which 50% inhibition is observed

Table 1. Select Confirmed GUS Inhibitors
Table 1. Contd.....

| Compound | Asinex Identifier a | Structure | IC₅₀ (µM) b |
|----------|---------------------|-----------|------------|
| 7        | BAS 02056251        | ![Structure Image] | 4.0        |
| 8        | ASN 03110025        | ![Structure Image] | 4.8        |

aChemical identifier number provided by Asinex Corporation
bFor IC₅₀ determinations, serial dilutions of compounds were tested starting at a high concentration of 20 µM. Average IC₅₀ values (n=3) are shown.

Fig. (8). IC₅₀ value determination for two screen hits. Compounds were serially diluted in 100% DMSO then transferred to assay plates for the GUS activity assay. Concentration response curves are shown for two representative hits from the screen.

DISCUSSION

GUS has been used as a reporter enzyme for many years. It is also the enzyme whose activity is detected in traditional tests for bacterial contamination in water samples. More recently, GUS appears to be responsible for converting the inactive CPT-11 metabolite back to active compound in the intestine. Active SN-38 in the intestine causes the dose-limiting diarrhea frequently associated with CPT-11 chemotherapy. These data have led to the hypothesis that small molecules that inhibit bacterial GUS activity may have potential therapeutic value in preventing CPT-11 mediated diarrhea. Limiting this side-effect may allow increased tolerance for the therapy and/or allow higher doses resulting in better therapeutic outcome.

In this report, we have adapted and validated a high throughput assay to identify inhibitors of GUS activity and used this assay to screen a diverse library of small molecules. The assay signal was shown to increase in a linear fashion with enzyme concentration and time. The assay was also tolerant up to at least 4% DMSO. As part of the validation, the control inhibitor D-Glucaric acid-1,4-lactone was used to determine IC₅₀ values using the final HTS assay with automation. The IC₅₀ values obtained were consistent between experiments, with the individual values within 12% of the mean. Whole 384-well plate variability studies using pre-spotted DMSO plates and automation were done to assess the variability of the assay for detection of inhibitors. The results generated Z'-factors of 0.81 for the inhibition assay window, indicating a robust assay for detection of hits.

As is typical for HTS validation, we screened a small set of compounds (the Prestwick collection) using the automated assay, to assess the performance of the assay in the presence of diverse compounds. This set of four plates resulted in Z'-factors of ≥0.8. An active rate of 3.6% was obtained which predicted in part, the high 5.1% active rate obtained from the Asinex library screen. The higher than expected active rates may be due the very low enzyme concentration in the assay (50 pM). With no other protein present and only minimal detergent present (0.01% Triton X-100), the assay may have been more susceptible to compounds that aggregate or have reactive impurities present. This explanation is supported by the observation that 72% of the confirmed actives had Hill slopes that were outside the 0.8 to 1.2 range. It has been reported that compounds that form aggregates in aqueous solu-
tion can non-specifically inhibit purified enzymes, usually generating steep Hill slopes (greater than 1.0) in IC50 value determinations [32, 33]. The higher active rate obtained in the screen compared to the Prestwick set may be, in part, due to highly related compounds represented in the Asinex library as suggested by the number of quinolinone thiourea compounds identified as hits. Lowering the compound screening concentration, for instance to 1 μM, or including bovine serum albumin (BSA) in the assay buffer are potential simple solutions that could reduce this hit rate.

From this library screen, at least 26 inhibitors of GUS were identified. The hits from Table 1 represent compounds that are significantly more potent than the control inhibitor. Five of the 8 inhibitors shown here are structurally novel inhibitors of GUS. One scaffold that emerged from the hit list was the quinolinone thiourea scaffold and highly related structures. Four of these compounds were the subject of published follow-up studies that demonstrated that these compounds inhibit GUS in living bacteria with IC50 values ranging from 18 to 1,300 nM without affecting cell viability [29]. They also did not inhibit mammalian GUS. One of these compounds was co-administered to mice receiving CPT-11 and the result that this compound significantly inhibited CPT-11-induced diarrhea. Thus, this GUS assay was successful in finding novel inhibitors of GUS activity for further study.

CONCLUSIONS

We have developed and validated a fluorescent high throughput assay for the discovery of small molecule inhibitors of GUS activity. Whole plate and IC50 variability studies confirmed the robustness of this assay for HTS. A chemical library was screened resulting in 26 confirmed inhibitors with good Hill slopes (0.8 – 1.2) with IC50 values ranging from 50 nM to 4.8 μM. These compounds may be used as starting points for the development of specific small molecule GUS inhibitors as chemical probes. Taken together, the GUS assay described herein is amenable for the discovery of inhibitors of bacterial GUS.

CONFLICT OF INTEREST

A method of use patent has been filed for the application of some of these compounds for CPT-11-induced toxicity.

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ABBREVIATIONS

GUS  = β-glucuronidase
DMSO  = dimethyl sulfoxide
RFU  = relative fluorescence units
GI  = gastrointestinal
4MUG  = 4-methylumbelliferyl glucuronide
E. coli  = Escherichia coli
IPTG  = isopropyl-1-thio-D-galactopyranoside

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