Combined Rational Design and a High Throughput Screening Platform for Identifying Chemical Inhibitors of a Ras-activating Enzyme*

The Ras family small GTPases regulate multiple cellular processes, including cell growth, survival, movement, and gene expression, and are intimately involved in cancer pathogenesis. Activation of these small GTPases is catalyzed by a special class of enzymes, termed guanine nucleotide exchange factors (GEFs). Herein, we developed a small molecule screening platform for identifying lead hits targeting a Ras GEF enzyme, SOS1. We employed an ensemble structure-based virtual screening approach in combination with a multiple tier high throughput experimental screen utilizing two complementary fluorescent guanine nucleotide exchange assays to identify small molecule inhibitors of GEF catalytic activity toward Ras. From a library of 350,000 compounds, we selected a set of 418 candidate compounds predicted to disrupt the GEF-Ras interaction, of which dual wavelength GDP dissociation and GTP-loading experimental screening identified two chemically distinct small molecule inhibitors. Subsequent biochemical validations indicate that they are capable of dose-dependently inhibiting GEF catalytic activity, binding to SOS1 with micromolar affinity, and disrupting GEF-Ras interaction. Mutagenesis studies in conjunction with structure-activity relationship studies mapped both compounds to different sites in the catalytic pocket, and both inhibited Ras signaling in cells. The unique screening platform established here for targeting Ras GEF enzymes could be broadly useful for identifying lead inhibitors for a variety of small GTPase-activating GEF reactions.

The Ras family of small guanosine triphosphatases (GTPases) is a large family of proteins consisting of over 20 members (1). This class of proteins regulates numerous cellular processes, such as cell growth, survival, movement, cytoskeletal organization, polarity, and gene expression (1, 2). Most Ras GTPases are regulated by two major classes of modulators, which control the cycling of these proteins between their inactive GDP-bound form and active GTP-bound form. The guanine nucleotide exchange factors (GEFs)2 catalyze the

2 The abbreviations used are: GEF, guanine nucleotide exchange factor; PH, pleckstrin homology; MEF, mouse embryonic fibroblast; PDB, Protein Data Bank; FL, fluorescein; TR, Texas Red; MTS, microscale thermophoresis; SAR, structure-activity relationship; EGFR, epidermal growth factor receptor; DH, Dbll homology domain; HTS, high throughput screening; uHTS, ultra-HTS; MTS, 3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium, inner salt.

Background: GEFs can serve as targets in Ras family GTPase activity signaling.

Results: A virtual screening coupled with the experimental screening platform is established targeting a Ras GEF enzyme. Conclusion: The multiple tier screening platform is useful for lead discovery targeting GEFs of Ras-like small GTPases. Significance: This study provides a unique screening approach for inhibitors of Ras and Ras-like GTPase activities.
could specifically suppress a Ras GEF (17). Here, we report an affordable academic drug screening platform for targeting potentially more druggable GEF enzymes, using the Ras GEF, SOS1, as a model study. We employed an ensemble structure-based docking approach in combination with a multiple tier high throughput experimental screening scheme utilizing two distinct, complementary fluorescent guanine nucleotide exchange assays, i.e. fluorescein-based GDP-dissociation assay and Texas Red-based GTP-loading assay. Applying this unique screening platform, we identified two chemically distinct small molecule inhibitors of SOS1 catalytic activity. Our study establishes an academically feasible drug screening platform that can easily be used to identify lead inhibitors of a variety of small GTPase GEFs.

EXPERIMENTAL PROCEDURES

Plasmids and Reagents—Guanosine 5'-triphosphate sodium salt solution, Tween 20, sodium deoxycholate, sodium chloride (NaCl), and Trizma® hydrochloride were purchased from Sigma. Magnesium chloride (MgCl₂) was purchased from Acros Organics (Gael, Belgium). Guanosine diphosphate (GDP), dimethyl sulfoxide (DMSO), and Nonidet P-40 alternative were purchased from Calbiochem®-EMD Millipore (Billerica, MA). Human epidermal growth factor (EGF) was purchased from Life Technologies, Inc.

Residues 1–182 of human Cdc42 were subcloned from a full-length human Cdc42/pET vector using the XhoI and HindIII restriction enzyme sites. Residues 1–166 of human H-Ras were subcloned from the full-length human H-Ras/pcDNA3.1 + mammalian expression vector (Missouri S&T cDNA Resource Center, Rolla, MO) into the bacterial pET15b vector using the XhoI and HindIII restriction enzyme sites. Residues 1–182 of human Cdc42 were subcloned from a full-length human Cdc42/pET vector into the pET15b vector using the XhoI and HindIII restriction enzyme sites. The Ras exchanger motif and Cdc25 catalytic domains (residues 564–1049) of human SOS1 (SOS1cat) were subcloned from the full-length human SOS1 MGC clone (accession number BC140215) (Life Technologies, Inc.) into the bacterial pET15b vector using the XhoI and HindIII restriction enzyme sites. Residues 1–182 of human Cdc42 were subcloned from a full-length human Cdc42/pET vector into the pET15b vector using the XhoI and HindIII restriction enzyme sites. The DH and PH catalytic domains (residues 1227–1571) of human Intersectin 1 (ITSN1DH-PH) were subcloned from a pGEX bacterial expression vector backbone into the bacterial expression vector pET15b (Novagen®-EMD Millipore) using the XhoI and HindIII restriction enzyme sites. The crystal structure of SOS1 in complex with Ras (bound at both the catalytic and allosteric sites, PDB code 1X2D) (19), as well as the unbound SOS1 structure (PDB code 1X2D) (19) were used to build alternative models of the SOS1 enzyme for docking simulations. Probis and CASTp were used to analyze structurally equivalent sites and overall similarity between the catalytic and allosteric Ras-binding interfaces within SOS1.

The University of Cincinnati–Drug Discovery Center’s collection of 350,000 compounds was used for virtual screening. Gesteiger partial charges were used for both the SOS1 enzyme and chemical compounds. A multiple step virtual screening approach was implemented, using increasingly stringent parameters and gradually more extensive sampling by increasing the number of energy evaluations (from 1.0 × 10⁶ to 1.0 × 10⁷), Genetic Algorithm runs (10 to 33) and population size (from 75 to 150), as discussed in detail previously (20). Upon completion of initial fast screening, 30,000 top candidate compounds with high estimates of binding affinities were retained, followed by re-scoring using improved sampling in the refinement stage. The top 3000 candidate compounds were again re-scored, using extensive sampling, and then clustered to avoid over-representation of some chemical classes.

As a result, two sets of hits were obtained, 315 for the catalytic site and 392 for the allosteric site, which were selected using both their predicted median inhibition constant and entropy of clustering of poses in multiple docking runs (excluding compounds with entropies larger than 0.4 to filter out likely nonspecific binders). An increasing level of overlap between top hits for these two sites (as tested by direct overlap of lists as well as clustering of the superset) was observed with subsequent runs and improved sampling, consistent with the observed structural similarity between the two sites. This was used as indication of the convergence of the sampling. Because of the overlap between the two sets, the set of hits selected for experimental validation consisted of 418 chemical compounds that were sufficient to cover the observed classes and clusters of hits.

Candidate compounds were further assessed from the point of view of their properties. Chemmine was used to cluster and visually analyze candidate molecules. Spipder and Polyview were used to analyze protein–protein complexes and to assess predicted docking poses for candidate inhibitors (the latter task facilitated by Polyview-MM) (21). Additionally, the predicted docking poses of candidate inhibitors were used to rationally guide which amino acid residues to mutate in SOS1 for site of action mutagenesis studies.

Using the BIOVIA Pipeline Pilot (version 8.5.0.200), computational structural similarity searches based upon both UC-773587 and UC-857993 were performed using the University of Cincinnati–Drug Discovery Center’s 350,000 compound library collection. As a result of these searches, 1 structurally similar compound to UC-773587 and 10 structurally similar compounds to UC-857993 were selected for experimental test-
ing. In addition, Chemmine was used to perform computational structural similarity searches for UC-773587 on a subset of 118,500 drug-like synthetic compounds from the NCI/DTP Open Chemical Repository (dtp.cancer.gov). As result of this search, seven structurally similar compounds to UC-773587 were selected for experimental testing.

**Protein Purifications**—Wild-type and mutant recombinant human SOS1cat (residues 564–1049) containing both the REM and Cdc25 domains, recombinant human ITSN1DH-PI (residues 1227–1571) containing both the DH and PH domains, recombinant human C-terminal truncated Cdc42 (residues 1–182), and recombinant human C-terminal truncated H-Ras (residues 1–166) were expressed in BL21(DE3) Escherichia coli as the N-terminal His6-tagged fusion proteins using the pET expression system (Novagen). For the wild-type and mutant recombinant human guanine nucleotide exchange factors (SOS1cat and ITSN1DH-PI), E. coli cells were grown to an 

\[ A_{600} \] between 0.4 and 0.8 and induced with 1 mM isopropyl \( \beta \)-D-1-thio-galactopyranoside overnight at room temperature. For the recombinant C-terminal truncated human GTPases (H-Ras and Cdc42), E. coli cells were grown to an \( A_{600} \) between 0.4 and 0.8 and induced with 1 mM isopropyl \( \beta \)-D-1-thio-galactopyranoside for 4 h at 37°C. Subsequently, the histidine-tagged fusion proteins were purified using Ni\(^{2+}\)-agarose affinity chromatography followed by gel filtration with a HiLoad 16/60 Superdex 200 column and FPLC AKTA Purifier UPC 10 system (GE Healthcare).

**Experimental High Throughput Guanine Nucleotide Exchange Assays**—For both the FL-based dissociation and the TR-based GTP loading, high throughput screening exchange assays, the Plate::Explorer\textsuperscript{TM} automated uHTS system (PerkinElmer Life Sciences) at the University of Cincinnati-Drug Discovery Center was employed to carry out the experimental screening scheme. For the FL-based dissociation assay, purified His\(_6\)-H-Ras (residues 1–166) was pre-loaded with BODIPY-FL-GDP for 1 h at room temperature with a stoichiometric ratio of 4:1, respectively. Exchange buffer (EB) (20 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 mM MgCl\(_2\), 0.01% Nonidet P-40 alternative, 1 mM DTT), 50 nM purified His\(_6\)-SOS1cat, 50-fold excess GTP, and DMSO or 100 \( \mu \)M chemical compounds were sequentially added to wells of a black 96-well plate (Corning Glass, catalog no. 3655) using 384-well automated liquid dispensers in conjunction with the Plate::Explorer\textsuperscript{TM} automated uHTS system (PerkinElmer Life Sciences). Subsequently, after a brief 30-s shake, the plate was incubated for 10 min at room temperature. Then, the kinetic reaction was initiated by adding 2 \( \mu \)M purified His\(_6\)-H-Ras (residues 1–166) immediately followed by 2.5-fold excess of BODIPY-TR-GTP with 384-well automated liquid dispensers in conjunction with the Plate::Explorer\textsuperscript{TM} automated system (PerkinElmer Life Sciences) (30 \( \mu \)l per well, final reaction volume). The kinetic reaction for both exchange assays was measured continuously for 15 min at room temperature in an EnVision plate reader (PerkinElmer Life Sciences) (fluorescein channel: excitation, 485 nm; emission, 535 nm; Texas Red channel: excitation, 560 nm; emission, 635 nm).

**96-Well Guanine Nucleotide Exchange Assays**—Initially, for the fluorescein-based dissociation exchange assay, purified His\(_6\)-H-Ras (residues 1–166) was pre-loaded with BODIPY-FL-GDP for 1 h at room temperature with a stoichiometric ratio of 4:1, respectively. For both the FL-based dissociation exchange assay, EB (20 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 mM MgCl\(_2\), 0.01% Nonidet P-40 alternative, 1 mM DTT), 50-fold excess of free GTP, and 50 nm purified His\(_6\)-SOS1cat were sequentially added to wells of a black 96-well plate (Corning Glass, catalog no. 3915). Subsequently, DMSO and the indicated concentrations of UC-773587 or UC-857993 were added to appropriate wells, followed by shaking the plate for 30 s and 10 min of incubation at room temperature. The reaction was initiated by adding 2 \( \mu \)M of pre-loaded H-Ras to each well (final reaction volume, 100 \( \mu \)l).

For the TR-based GT loading exchange assay, EB and 100 nm purified His\(_6\)-SOS1cat were sequentially added to the wells of a black 96-well plate. Subsequently, DMSO and the indicated concentrations of UC-773587 and UC-857993 were added to the appropriate wells, followed by shaking the plate for 30 s and 10 min of incubation at room temperature. The reaction was initiated by adding 2 \( \mu \)M of pre-loaded H-Ras to each well (final reaction volume, 100 \( \mu \)l). The kinetic reactions for both exchange assays were measured continuously for 15 min at room temperature in an EnVision plate reader (PerkinElmer Life Sciences) (fluorescein channel: excitation, 485 nm; emission, 535 nm; Texas Red channel: excitation, 560 nm; emission, 635 nm).

**Microscale Thermophoresis (MST) Binding Assays**—Purified wild-type and mutant His\(_6\)-SOS1cat and C-terminally truncated His\(_6\)-H-Ras were labeled with the red NT-647 amine-reactive dye using a protein labeling kit (NanoTemper Technologies). The indicated concentrations of UC-773587 or UC-857993 were incubated with NT-647-labeled SOS1cat (100 nm) or H-Ras (100 nm) in binding buffer (20 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 mM MgCl\(_2\), 0.1% Triton X-100, and 2% DMSO) at room temperature in the dark for 30 min. For the competition of binding experiments, purified His\(_6\)-SOS1cat was labeled with the red NT-647 cysteine-reactive dye using a protein labeling kit (NanoTemper Technologies). 0.2 to 100 \( \mu \)M unlabeled purified C-terminally trun-
cated His\textsubscript{6}-H-Ras were incubated with NT-647-labeled SOS1-cat (50 nM) in the presence of the indicated concentrations of UC-773587 or UC-857993 in binding buffer (20 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 mM MgCl\textsubscript{2}, 0.1% Tween 20, and 2% DMSO) at room temperature in the dark for 30 min. Subsequently, samples were loaded into hydrophilic capillaries (NanoTemper Technologies), and binding events were measured with a Monolith NT.115 reader (NanoTemper Technologies). Binding data were analyzed using Thermophoresis or Thermophoresis with Tempera-
Antibodies were diluted (1:1000) in TBS-T buffer containing 5% BSA. Membranes were developed by ECL Western blot detection reagents (GE Healthcare and Thermo Fisher Scientific). Images for the ECL-developed membranes were taken by the Fuji Image Reader LAS1000 or ChemiDoc Touch Imaging System (Bio-Rad).

### RESULTS

**Combined Virtual Screening and High Throughput Experimental Screening to Identify Inhibitors of GEF Catalytic Activity**

Multiple crystal structures of the Ras GEF, SOS1, and its complexes (PDB code 1XD2 and 1XD4) (Fig. 1A) were used for virtual screening to guide an ensemble docking approach targeting both the catalytic and allosteric sites within SOS1 to identify candidate inhibitors (19). Detailed structural analysis of the SOS1-Ras complexes revealed structural similarity between Ras binding interfaces. In particular, similar conformations of the switch I region of Ras are involved in the interactions with SOS1 at both the catalytic and allosteric sites (Fig. 1B) were targeted in virtual screening. Utilizing ensemble docking with multiple (bound versus unbound) structures and comparing predicted docking poses at structurally equivalent docking sites allowed us to improve sensitivity, while reducing false-positive rates. In addition, integrating multiple structures into the virtual screening protocol allowed us to partially alleviate the limitation of rigid body docking.
FIGURE 2. High throughput screening platform for small molecule inhibitors of GEF catalytic activity. A, schematic of the methodology of the two complementary fluorescent (fluorescein and Texas Red) intensity guanine nucleotide exchange assays used for experimental screening. B, characterization of the BODIPY-FL-GDP dissociation primary screening assay. 50 nM purified His6-SOS1cat (red circle)-stimulated BODIPY-FL-GDP nucleotide exchange for unlabelled GTP on purified 2 μM His6-H-Ras (residues 1–166) (blue circle). C, validation of the GDP dissociation primary screening assay. −SOS1cat (blue circle) and +SOS1cat (red circle) controls from the primary screening assay plates are plotted using a 15-min time point and subsequently used to calculate the statistical coefficient, Z-factor (22), to validate the assay for HTS. D, HTS results of the primary screen. −SOS1cat (blue circle) and +SOS1cat (red circle) DMSO controls along with the chemical compounds (green circle) tested for their ability to inhibit SOS1 catalytic activity are plotted using a 15-min time point. The >30% inhibition cutoff used is indicated. E, HTS platform experimental screening scheme used to identify inhibitors of SOS1 catalytic activity. 418 candidate compounds were identified from the virtual screen as described in Fig. 1. 394 of these 418 compounds were available for experimental screening. 23 of the 394 compounds were identified as hits from the primary screen. 15 of these 23 hits were confirmed using two complementary fluorescent intensity assays. Of these 15 hits, 10 hits were validated by a 10-point dose response (2-fold serial dilution; 0.195–100 μM), and subsequently 9 of these 10 hits were validated in 96-well format dose response (0, 2, 5, 10, 20, 50, and 100 μM). Five of the nine hits showed activity in both complementary fluorescent assays. Two of these five hits were selected for further validation and characterization. Data in B–D are expressed as percent change of fluorescence units normalized to the initial time point. Data in B were performed in triplicate. C, three outliers were removed from both the −SOS1cat and +SOS1cat control datasets based upon visual inspection.
Beginning with the University of Cincinnati-Drug Discovery Center’s 350,000 compound collection, a multistage virtual screening protocol was used to identify the top 30,000 hits with limited sampling. Subsequently, using better sampling, the 30,000 hits were reduced to 3000 top hits (Fig. 1C) (20). At each stage, alternative conformers of Ras interaction at both the cat-

**FIGURE 3. Representative hit validation in 96-well plates.** Dose-dependent inhibition of 50 nm purified His<sub>6</sub>-SOS1<sub>cat</sub> (black circle)-mediated GDP for GTP nucleotide exchange upon 2 μM His<sub>6</sub>-H-Ras (residues 1–166) (gray circle) in both 96-well BODIPY-FL-GDP dissociation and BODIPY-TR-GTP loading assays by the five validated hit compounds that displayed activity in both complementary experiments. Chemical structures of the five hit compounds are displayed on the left side of their respective experimental data. Data are expressed as percent change of fluorescence units normalized to the initial time point. Data were performed in triplicate and represent the mean.
Analytic and allosteric sites on SOS1 were used to perform docking simulations to partially account for conformational changes related to SOS1-Ras complex formation and to provide consistency-based filtering along with improved ranking of candidate inhibitors. Top hits for each site on SOS1, rank base on high predicted binding affinity and consistent binding to a specific FIGURE 4. UC-773587 and UC-857993 inhibit GEF catalytic activity. A, chemical structures of the two hit compounds, UC-773587 and UC-857993. UC-773587 and UC-857993 dose-dependently inhibit (B and D) 50 nM His<sub>6</sub>-SOS1<sub>cat</sub> (black circle)-mediated BODIPY-FL-GDP for GTP nucleotide exchange on 2 μM His<sub>6</sub>-H-Ras (residues 1–166) (gray circle) or (C and E) 50 nM His<sub>6</sub>-SOS1<sub>cat</sub> (black circle)-mediated BODIPY-TR-GTP loading on 2 μM His<sub>6</sub>-H-Ras (residues 1–166) (gray circle) in a 96-well format at the indicated compound concentrations. F, UC-773587 (3 μM) and UC-857993 (20 μM) additively inhibit 50 nM His<sub>6</sub>-SOS1<sub>cat</sub> (black circle)-mediated BODIPY-FL-GDP for GTP nucleotide exchange on 2 μM His<sub>6</sub>-H-Ras (residues 1–166) (gray circle) in a 96-well format. Data in B–F are expressed as percent change in fluorescence units normalized to the initial time point and were performed in triplicate. Data in B–E represent the mean of three independent experiments.
site in a dominant pose within the simulation box, as captured by low entropy of clustering poses acquired in multiple docking runs (Fig. 1D), were combined and clustered based upon chemical similarity. This resulted in a set of 418 candidate compounds, of which 394 compounds were available, for experimental screening in our high throughput screening platform (Figs. 1D and 2E).

For our high throughput screening platform, two complementary fluorescent intensity guanine nucleotide exchange assays were developed (28–30). One assay utilized the BODIPY-FL-GDP nucleotide to measure SOS1\textsubscript{cat}-mediated GDP dissociation on H-Ras that results in a loss of fluorescence, and the other assay utilized a BODIPY-TR-GTP nucleotide to measure SOS1\textsubscript{cat}-mediated GTP loading to H-Ras resulting in an increase in TR fluorescence (Fig. 2A). Integrating these two complementary assays into our high throughput screening platform scheme enabled us to identify inhibitors of SOS1 catalysis that mechanistically could disrupt two distinct steps of the GTPase cycle. Additionally, using two different fluorescent channels (i.e. fluorescein and Texas Red) as a read-out for each complementary guanine nucleotide exchange assay reduced the possibility of identifying false-positives as a result of fluorescent artifact. Because of the reproducibility and robustness of the BODIPY-FL-GDP dissociation guanine nucleotide exchange assay, it was selected to perform the primary screening stage of the multiple tier high throughput screening platform scheme (Fig. 2E). Using a time point of 15 min, the fluorescein-based dissociation assay produced a combined $Z_\text{H}\text{.11003}/Z_\text{H}\text{.11004}/Z_\text{H}\text{.11005}/Z_\text{H}\text{.11006}$ factor of 0.5 using the positive and negative controls (−SOS1\textsubscript{cat} and +SOS1\textsubscript{cat}) from each primary screening assay plate for calculation, indicating it to be an excellent assay for primary high throughput screening (Fig. 2B, D and E). Using the dissociation assay, 394 candidate compounds at a dose of 100 $\mu M$ were tested for their ability to inhibit SOS1\textsubscript{cat} catalytic activity. Using a cutoff of >30% inhibition, 23 hits were identified for a 5.8% hit rate (Fig. 2D and E). Typically, one would expect a hit rate of ≤1% for high throughput screens using diverse chemical libraries (28, 31). However, our computer-aided virtual screen
was able to enhance the screening hit rate to 5.8%, thus indicating the efficiency of combining computer-guided virtual screens with experimental high throughput screening platform campaigns. Subsequently, the 23 hit compounds at a dose of 100 μM were tested for their ability to inhibit SOS1 catalytic activity using a cutoff of >30% inhibition in both the dissociation and GTP loading assays. 15 of the 23 hit compounds were confirmed as true hits reducing the hit rate to 3.8%. Nine of these 15 hit compounds were active in the dissociation assay, and six of the 15 hit compounds were active in both the dissociation and GTP loading assays. These 15 hit compounds were further tested for their ability to inhibit SOS1cat catalytic activity in a 10-point dose-response assay using both fluorescent intensity guanine nucleotide exchange assays. 10 of the 15 hit compounds dose-dependently inhibited SOS1cat catalytic activity further reducing the overall hit rate to 2.5%. Of these 10 compounds, five compounds were only active in the dissociation assay, and the remaining five compounds were active in both the dissociation and GTP loading assays. At this stage, fresh powder of the 10 hit compounds was obtained and manually tested for their ability to dose-dependently inhibit SOS1cat catalytic activity in both the dissociation and GTP loading assays (Fig. 3). Two of the five validated compounds were selected for follow-up characterization due to their experimental efficacy and selectivity. Three of the five validated compounds were ruled out due to their lack of selectivity (UC-773587) for direct binding to SOS1 versus H-Ras, poor efficacy (UC-104460), and/or poor solubility (UC-992652).

**Biochemical Characterization of Two Chemically Distinct Inhibitors of SOS1 Catalytic Activity**—Two chemically distinct inhibitors (Fig. 4A) of SOS1 catalytic activity were further characterized as inhibitors of SOS1 in a series of follow-up biochemical experiments. First, UC-773587 and UC-857993 were tested for their ability to inhibit SOS1cat catalytic activity in our two complementary fluorescent intensity guanine nucleotide exchange assays. Both UC-773587 and UC-857993 dose-dependently inhibited SOS1cat-mediated GDP dissociation and GTP loading on H-Ras (Fig. 4, B–E). Interestingly, combination treatment of UC-773587 and UC-857993 additively disrupted the interaction between SOS1cat and GDP-bound H-Ras (Fig. 5G). To confirm that UC-773587 and UC-857993 inhibited both SOS1 catalytic activity and the SOS1-Ras interaction by binding to the expected target SOS1 and not H-Ras, direct binding experiments to SOS1 and H-Ras were carried out using microscale thermophoresis. As predicted, UC-773587 and UC-857993 were found to directly bind to SOS1cat (UC-773587 - Kd = 3.4 μM; UC-857993 - Kd = 14.7 μM) but not H-Ras with micromolar affinity (Fig. 5A–D). Also, consistent with our virtual screening model, UC-773587 and UC-857993 dose-dependently disrupted the SOS1-Ras interaction by competitively inhibiting the binding of SOS1cat to GDP-bound H-Ras in a microscale thermophoresis assay (Fig. 5, E and F).
Third, to test whether our compounds were selective for the Ras GEF, SOS1, versus the distantly related Rho GEF, ITSN1, UC-773587, and UC-857993, was tested for the ability to inhibit SOS1cat-mediated and the DH-PH catalytic module of Intersectin 1 (ITSN1DH-PH)-mediated GDP nucleotide exchange on H-Ras and Cdc42, respectively. Interestingly, both UC-773587 and UC-857993 showed selective inhibition of SOS1cat versus ITSN1DH-PH catalytic activity (Fig. 6). Given the fluorescent basis of these biochemical experiments, UC-773587 and UC-857993 were tested for their UV-visible absorbance spectrum and/or emission spectrum in the fluorescein and Texas Red channels to rule out spectral interference. Both compounds displayed no absorbance peaks at the wavelengths used for the biochemical experiments, and furthermore, UC-773587 did not show autofluorescence in either the fluorescein or Texas Red channels (Fig. 7).

Mapping the Sites of Action for UC-773587 and UC-857993—Based on our docking models, strong efficacy in our GEF exchange assays, additive inhibition of UC-773587 and UC-857993 in our biochemical experiments, and the observation that UC-773587 and UC-857993 disrupt the interaction between SOS1 and GDP-bound H-Ras, we hypothesized that UC-773587 and UC-857993 mapped to the catalytic site on SOS1. Therefore, we performed alanine scanning mutagenesis studies of residues predicted to interact with our compounds at the catalytic site (Fig. 8, A and B) by generating 19 single alanine mutants and five double alanine mutants. Additionally, we generated a single alanine mutant at the allosteric site (L687A), which has been shown previously to be critical for Ras binding (19, 32), as a negative control. Of the single point mutants, two (K939A and H911A) completely disrupted binding and one mutant (L934A) enhanced binding of UC-857993 to SOS1 (Fig. 8, C and D).
9, B–D and F). Conversely, for UC-773587, two single point mutants (I825A and K814A) reduced binding and the double mutant (I825A/K814A) completely disrupted binding to SOS1, and the single point mutant of a residue at the bottom of the predicted binding pocket (M824A) slightly enhanced binding (Fig. 9, A and C–E). For both compounds, mutating one of the critical residues, Leu-687, at the allosteric site to alanine did not significantly affect compound binding (Fig. 9, A and B). The lack of binding, reduced binding, or enhanced binding was unlikely due to improper protein folding as four of the mutants (K814A, M824A, I825A, and K939A) were catalytically active, although the three other mutants (H911A, L934A, and K814A/I825A) were catalytically dead (Fig. 10), but the inactive mutants have been shown previously to be critical for the SOS1-Ras interaction (33, 34). Interestingly, it has been shown that His-911 and Lys-939 residues of SOS1 are critical for the interaction of SOS1 with the switch I region of Ras, whereas the Ile-825 residue of SOS1 is critical for the interaction of SOS1 with the switch II region of Ras (34). Based upon the lack of binding of UC-857993 to the H911A and K939A mutants, UC-857993 maps to the switch I Ras interaction region at the catalytic site of SOS1. The enhanced binding of UC-857993 to the L934A mutant and UC-773587 to the M824A mutant is likely due to removal of steric hindrance allowing for a better binding pocket. Based upon reduced binding of UC-773587 to the I825A single mutant and complete inhibition of binding to the K814A/I825A double mutant, UC-773587 may map to the switch II Ras interaction region at the catalytic site of SOS1. The reduced binding of UC-773587 to the K814A single mutant and subsequent complete inhibition of binding to the K814A/I825A double mutant may be attributed to an indirect conformational effect coupling the two mutated sites.

**SAR Studies for UC-773587 and UC-857993**—To further understand the structure-activity relationship of our two inhib-
itors of SOS1, computational searches for structural analogs of UC-773587 and UC-857993 were undertaken. Subsequently, the structural analogs were tested for their ability to inhibit SOS1_{cat}-mediated GDP nucleotide exchange on H-Ras. Limited availability of closely related analogs allowed only limited SAR conclusions. A carboxyl function appears to be essential because all of the activities (IC_{50}/H11021\textsuperscript{50}/H9262 M) contain this group, most clearly demonstrated by the loss of activity from the carboxyl function.
boxylate NSC-76322 to its methyl ester NSC-76323. H-bonding
groups present at the distal ends of the molecule (as in
UC-773587) also improve potency (Fig. 11). Interestingly,
appending the carboxylate via a linkage that can adopt a per-
pendicular conformation (e.g. UC-773587 and NSC-76322) to
the polycyclic system affords greater activity than the direct
appending of the carboxylate as in NSC-9608.

For UC-857993, analog availability allowed a more targeted
range of modifications to the UC-857993 scaffold to be
explored. Modification of the annulated cyclopentene ring
modulated activity in the following manner: (i) deletion of the
ring as in UC-384876 and UC-415564 eliminated all activity; (ii)
expansion of the ring to a six-membered saturated ring as in
UC-397847 led to a slight loss in activity; but (iii) replacing the
ring with a flexible \( n \)-pentyl group (UC-397299) increased activity 2-fold. The appended aryl system is essential for activity (note: the loss of activity in analogs UC-501950 and UC-404796). Minor changes in substitutions as in UC-917210 and UC-781482 show little effect on activity. The carboxylate function, which most likely interacts with the Lys-939 residue in SOS1, is critical as shown by the complete loss of activity when converted to a methyl ester as in UC-503495 or replacement with bromine in UC-782008 (Fig. 12).

**UC-773587 Structure-Activity Relationship (SAR)**

![Diagram of UC-773587 and related structural analogs](image)

| Compound #   | IC50 (\( \mu \)M) | SE (\( \mu \)M) |
|--------------|-------------------|---------------|
| UC-773587    | 4.5               | 1.20          |
| NSC-126404   | NA                | NA            |
| NSC-76322    | 15.3              | 1.18          |
| NSC-76323    | NA                | NA            |
| NSC-95170    | NA                | NA            |
| NSC-9608     | 29.1              | 1.13          |
| UC-387626    | NA                | NA            |

(NA = No Activity)

![Figure 11. Structure-activity relationship for UC-773587.](image)

**UC-773587 and UC-857993 Inhibit Ras Signaling and Cancer Cell Growth**—To assess cellular activity, the two SOS1 inhibitors (UC-773587 and UC-857993) were tested in cells at inhibiting epidermal growth factor receptor (EGFR)-mediated Ras activation and downstream ERK activation (35, 36). Therefore, we serum-starved NIH/3T3 cells and pretreated them with either UC-773587 or UC-857993. Subsequently, we stimulated the cells with EGF to initiate Ras signaling. Both UC-773587 and UC-857993 dose-dependently inhibited (<50% inhibition) growth of NIH/3T3 cells (Fig. 13, A and B). Both UC-773587 and UC-857993 dose-dependently inhibited Ras activation and downstream ERK activation mediated through EGFR-SOS1-Ras-Raf1-MEK-ERK signaling (Fig. 13, C and D). Furthermore, to test the selectivity of both compounds for SOS1-Ras signaling, a GTPase-activating protein activity-deficient H-Ras(G12V) mutant expressing the mouse embryonic fibroblast cell line was produced (3, 25). Therefore, we tested the ability of UC-773587 and UC-857993 to inhibit growth of wild-type mouse embryonic fibroblasts versus activated mutant H-Ras(G12V)-transformed mouse embryonic fibroblasts. Interestingly, under the experimental conditions tested, UC-773587 and UC-857993 specifically inhibited wild-type mouse embryonic fibroblast growth but not the growth of oncogenic H-Ras(G12V)-expressing mouse embryonic fibroblasts (Fig. 13, E and F). As a result,
this suggests that the compounds characterized here are selective for SOS1-Ras signaling.

To further explore the selectivity of these inhibitors for SOS1-Ras signaling in a more translational model, UC-773587 and UC-857993 were tested for their effect upon cell growth on a panel of human cancer cell lines with wild type or activated mutant Ras status. Under the experimental conditions tested, both compounds specifically inhibited growth of prostate (DU-145) and pancreatic (BxPC-3) cancer cells containing the wild-type Ras status but not that of pancreatic (Panc-1) cancer cells containing the activated mutant K-Ras status (Fig. 14, A and B). Additionally, UC-773587 and UC-857993 inhibited downstream ERK activation in DU-145 prostate cancer cells with the wild-type Ras status but not in Panc-1 pancreatic cancer cells with the activated mutant K-Ras status (Fig. 14, C and D). Therefore, consistent with the mouse embryonic fibroblast results, these human cancer cell line results further suggest that the UC-773587 and UC-857993 are selective for SOS1-Ras signaling. Interestingly, there was an exception in which cell growth of MIA-PaCa-2 pancreatic cancer cells containing activated mutant K-Ras was suppressed by both UC-773587 and UC-857993 (Fig. 14, A and B), but these cells have previously been shown to be dependent upon SOS1 for their growth through a unique autocrine loop dependent upon wild-type H- and N-Ras (37, 38).

It has been shown previously that SOS1 is differentially expressed in prostate and breast cancer populations (7, 8) and is required for Ras signaling and growth of human prostate cancer cells (7). Given the additive effects of combination treatment of UC-773587 and UC-857993 in our biochemical experiments, we used the SOS1-dependent human DU-145 prostate cancer cells as a model to examine whether combination treatment of our SOS1 inhibitors could additively inhibit growth of the DU-145 cancer cells. As expected, both UC-773587 and UC-857993 were able to individually inhibit DU-145 cell growth in a dose-dependent manner (Fig. 14, E and F). From this result, modest inhibitory doses for UC-773587 and UC-857993 were selected to assess additivity of the compounds in the DU-145 cancer cell growth model. Consistent with the in vitro biochemical results, combination treatment of UC-773587 and UC-857993 inhibited growth of the DU-145 cells (Fig. 14G). Taken together, these cellular results suggest that both SOS1 inhibitors map to the catalytic site on SOS1 but can work additively or synergistically in suppressing SOS1 activity. 
DISCUSSION

Currently, available pharmacological tools to dissect small GTPase signaling mechanisms remain limited. Although there has been significant efforts in targeting post-translational modifications (i.e. farnesyltransferases and geranylgeranyltransferases) of small GTPases (39–43) and small GTPase effector proteins (e.g. Raf, PI3K, ROCK, and PAK) (1, 44), these lead chemical inhibitors lack selectivity for upstream GTPase signaling. We and others have attempted to directly target small GTPases (i.e. Rac1, RhoA, Cdc42, and Ras) themselves (14–16, 45–49), but these chemical inhibitors tend to be of low affinity/efficacy due to the difficulty of targeting globular GTPase structures. With the exception of the Rho GEF inhibitors ITX3 for Trio and Y16 for LARG and the recently identified SOS1 inhibitor and activator (17, 50), there have been no other published reports successfully targeting the enzymatic activators of small GTPases, GEFs. The potentially structurally and functionally druggable GEFs are attractive targets for inhibition as they link upstream signaling to small GTPases and subsequent cellular signaling. Here, we present an affordable and efficient screening platform combining structure-based virtual screening and an experimental high throughput screening platform in designing and discovery of GEF inhibitors and an application to the Ras GEF, SOS1. Given the conserved mechanism of GEF action,
we expect this screening platform can readily be adapted to studies of other small GTPase GEFs with available structural information.

Using our combinatorial screening platform, we were able to identify two chemically distinct inhibitors (UC-773587 and UC-857993) of the catalytic activity of SOS1. As predicted by FIGURE 14.

**FIGURE 14.** UC-773587 and UC-857993 inhibit Ras signaling and growth of pancreatic and prostate cancer cells. UC-773587 (A) and UC-857993 (B) were tested for their ability to inhibit a 3-day cell growth of a panel of pancreatic and prostate cancer cell lines. Cells were grown in the presence of DMSO or indicated concentrations of UC-773587 or UC-857993 for 3 days and subsequently assayed for their growth by MTS as described under “Experimental Procedures.” C and D, DU-145 cells and Panc-1 cells were serum-starved for 24 h and pretreated with indicated concentrations of UC-773587 or UC-857993 for 16 h (DU-145-UC-773587), 2 h (DU-145-UC-857993), or 1 h (Panc-1-UC-773587 and UC-857993). Activation of phospho-ERK was measured by Western blot. Cell growth of DU-145 cells was measured in the presence of DMSO or indicated concentrations of UC-773587 (E) or UC-857993 (F) individually or in combination (G, left panel) over 72 h. Every 24 h of growth of the cells was assayed by MTS. G, right panel, combination treatment growth inhibition is represented as a bar graph at 72 h to assess additivity. Cell growth data were performed in triplicate or quadruplicate and represent the mean ± S.E. Data in A and B are plotted as relative growth on day 3 normalized to the DMSO control. Western blot data are representative of n = 3 experiments.
our virtual screen, both inhibitors could bind directly to SOS1 and inhibited two distinct mechanistic steps of the GTPase cycle, including GEF-mediated GDP dissociation and GTP loading by interfering with the interaction between GDP-bound Ras and SOS1. Mutation analysis mapped both compounds to the catalytic pocket of SOS1 with UC-773587 binding to the Ras switch II interaction region and UC-857993 binding to the Ras switch I interaction region of SOS1. Interestingly, combination treatment additively inhibited SOS1 catalysis, blocked the SOS1-Ras interaction, and inhibited SOS1-dependent human prostate cancer cell growth. This suggests a potential new paradigm for dual targeting of catalytic pockets on enzymes with relatively large drug-accessible surfaces. In addition, recent studies have implicated SOS1 in oncogenic K-Ras tumorigenesis in pancreatic, colon, and breast cancer models by mediating H-Ras and N-Ras signaling through an autocrine loop (37, 38, 51), suggesting an interesting future utility of SOS1 inhibitors in K-Ras-mediated oncogenicity, which is supported here by the inhibition of MIA-PaCa-2 pancreatic cancer cell growth with both SOS1 inhibitors.

A recent review article (52) discussed the potential pitfalls of pan-assay interference compounds that often show up as false-positive hit compounds and trick chemical biologists into believing that they are true hits. Given the fluorescein-like structures of the hit compounds characterized here, we were concerned that they could be pan-assay interference compounds. To this end, we performed a series of careful validations and control experiments to confirm that this was not the case. First, in our experimental high throughput screening platform, we have built in a control for potential fluorescent artifact compounds by following up on hits using two complementary fluorescent-based biochemical exchange assays in both the fluorescein and Texas Red channels. Our hit compounds were validated to work in both exchange assays and fluorescence channels. Additionally, both compounds did not absorb light in the UV-visible spectra at wavelengths used for our biochemical experiments, and UC-773587 did not auto-fluoresce in the fluorescein or Texas Red channels, ruling out potential artifacts due to compound intrinsic fluorescence. Second, as suggested by the Baell and Walters (52), we performed follow-up validations with fresh powder compounds from our own source along with commercial sources to ensure the purity of our compounds and to show that they can directly bind to the target protein, SOS1, competitively disrupt the SOS1-Ras interaction, and selectively inhibit SOS1 catalytic activity specifically.

Third, mutagenesis studies show that the hit compounds do not covalently bind to the target protein and do not bind to several mutants, and a preliminary SAR analysis shows several chemical structural analogs were inactive. Finally, a cellular test shows that the hits are capable of dose-dependently inhibiting EGFR-SOS1-Raf1-MEF-ERK signaling and proliferation of SOS1-addicted prostate and pancreatic cancer cells. Despite these collective efforts, there are limitations to our biochemical studies of the hit compounds; more detailed structural work by crystallization of the hits bound to SOS1 would help guide future medicinal chemical development. Overall, these studies provide a proof-of-concept for multiple platform screening for inhibitors of a Ras GEF, and we expect the approach can be adapted and applied to identifying lead inhibitors of other GEF proteins in academically feasible laboratories.

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