An Integrated Computational and Experimental Binding Study Identifies the DNA Binding Domain as the Putative Binding Site of Novel Pyrimidinetrione Signal Transducer and Activator of Transcription 3 (STAT3) Inhibitors

Shan Sun1, Peibin Yue2, Mingzhu He3, Xiaolei Zhang3, David Paladino3, Yousef Al-Abed3, James Turkson4* and John K Buolamwini1,4**

1Department of Pharmaceutical Sciences, College of Pharmacy, University of Tennessee Health Science Center, 847 Monroe Avenue, Suite 327, Memphis, TN 38163, USA
2University of Hawaii Cancer Center, Cancer Biology and Natural Products and Experimental Therapeutic Programs, 701 Iilau Street, Suite 344, Honolulu, HI 96813, USA
3Center for Molecular Innovation, The Feinstein Institute for Medical Research, 350 Community Drive, Manhasset, NY 11030, USA
4Department of Pharmaceutical Sciences, College of Pharmacy, Rosalind Franklin University of Medicine and Science, 333 Green Bay Road, North Chicago, IL 60064, USA

Abstract

Objective: Signal transducer and activator of transcription 3 (STAT3) is as a potential drug target for cancer and inflammation. Strategies to block STAT3 dimerization have dominated STAT3 inhibitor discovery, while inhibition of direct STAT3-DNA binding has not been much explored. This study was to identify novel STAT3 inhibitors that may be developed into probes or therapeutics, and to investigate their putative binding site.

Methods: A library of in-house compounds was screened against STAT3-DNA binding using an electrophoretic mobility shift assay (EMSA). Inhibition of DNA and STAT3:STAT3 or STAT3:STAT1 interaction and antiproliferative activity in STAT3 expressing or STAT3 knockout cells were tested. Pharmacophore modeling and 3D-QSAR; and docking with molecular mechanics generalized born surface area (MM-GBSA) refinement were also undertaken at the STAT3 SH2 and STAT3 DNA binding domains. Surface plasmon resonance (SPR) analysis was done to determine STAT3 domain interactions. Organic synthesis was also undertaken.

Results: Pyrimidinetriones derivatives were identified as novel STAT3 inhibitors, with activity in low micromolar concentrations. Application of 3D-QSAR and docking analysis with MM-GBSA refinement suggested that the compounds bind at the STAT3-DNA binding interface, and not at the SH2 domain where most current STAT3 inhibitors are thought to bind. The results were confirmed with SPR analysis, pointing to the DNA binding domain (DBD) as the putative binding site of this novel class of STAT3 inhibitors. The analysis guided the synthesis of active novel compounds.

Conclusion: Pyrimidinetrione were identified as new STAT3 inhibitors that putatively bind at the STAT3 DBD. This study is an excellent example of the use of QSAR and structure-based design to aid the identification of putative ligand binding sites on proteins. The compounds will provide new tools for studying STAT3 biology, and also serve as potential leads for the development of new therapeutics against STAT3 to fight or study diseases such as cancer and inflammation.

Keywords: Docking; MM-GBSA; Pharmacophore-based 3D-QSAR; STAT3 DNA binding site; STAT3 inhibitors; Electrophoretic mobility assay; Surface plasmon resonance assay

Introduction

Signal transducers and activators of transcription (STATs) regulate gene expression in normal cellular responses to cytokines and growth factors [1-3]. STAT3 is the most studied one of the 7 family members identified due to its major mediating effects on carcinogenesis and inflammation [2,4]. The abberant activation of STAT3 has been detected in a wide variety of human cancer cell lines and tissues [5].

Constitutively active STAT3 suppresses apoptosis [6], upregulates the expression of cell proliferation genes [7,8], stimulates tumor angiogenesis [9] and opposes anti-tumor immune responses [10,11]. Hence, STAT3 is considered as a promising therapeutic target for cancer therapy.

STAT3 activation begins with the phosphorylation of a critical tyrosine residue (Tyr705) on its Src homology 2 (SH2) domain by activated growth factor receptors, Janus kinases (JAKs) or Src tyrosine kinase. Upon activation, STAT3 forms dimers through a reciprocal tyrosine residue (Tyr705) on its Src homology 2 (SH2) domain by activated growth factor receptors, Janus kinases (JAKs) or Src tyrosine kinase. Upon activation, STAT3 forms dimers through a reciprocal SH2 domain interaction and translocates to the nucleus where the dimers bind to the promoters of target genes and activate specific gene expression. For many years, the SH2 domain-pTyr dimerization step was regarded as an attractive target for interfering with STAT3 function and this strategy has been exploited in many STAT3 drug discovery approaches [12]. Most nonpeptide STAT3 inhibitors known to date are small molecule compounds and...
peptidomimetics, such as Stattic [13], STA-21 [14], S31-201 [15], LLL-12 [16], 14aa [17], etc., and are mainly STAT3 dimerization disrupting agents. Unfortunately, none of these STAT3 dimerization disrupting agents has reached the clinic as cancer therapeutics. In contrast to STAT3 dimerization disrupting agents, not much progress has been made in approaches aimed at direct inhibition of STAT3 DNA binding [18,19]. Recent studies have indicated that STAT3 nuclear import can take place constitutively and independently of Tyr705 phosphorylation through the nuclear import carrier importin-α3 [20]. In addition, the binding of unphosphorylated STAT3 directly to DNA has also been observed by an electrophoretic mobility shift assay (EMSA) [21]. These studies suggest that the strategy of blocking STAT3 dimerization by targeting the SH2 domain might be unable to effectively inhibit STAT3 activity. Although a number of molecules have been screened or developed as putative STAT3 SH2 domain binding inhibitors, binding sites for these small molecules at the SH2 domain have not been identified by any crystal structure.

In early 2000, a tetrahydrosobenzofurane derivative [22] and a class of platinum (IV) compounds [23,24] were reported to potentially interact with the STAT3 DNA binding domain (DBD) and in turn inhibited STAT3-DNA binding and transcriptional activities; but the exact mode of inhibition of STAT3 is not understood. A peptide aptamer has also been identified as a STAT3 inhibitor that specifically interacts with the STAT3 DBD [25,26]. Recently, selective STAT3 DBD inhibitors with the 5-phenyl-1H-pyrrolo-2(3H)-ketone core structure were reported to directly bind to the DBD and inhibit the DNA binding activity of STAT3 both in vitro and in situ [19,27]. The reason for the low interest in STAT3-DNA-binding interface could be the notion that DBDs of transcription factors are generally “undruggable”, often being flat and of limited selectivity [28,29]. The inability of any of the many small molecule STAT3 SH2 domain to progress to the clinic so far, gives cause to pursue other STAT3 domain binding inhibitors such as the DBD inhibitors.

We carried out an unbiased screening of in-house compound libraries using electrophoretic mobility shift assay (EMSA) to identify novel inhibitors preventing STAT3-DNA binding. The hits indicated that the pyrimidinetrione system constituted a novel STAT3 inhibitor scaffold. We further pursued it by testing additional pyrimidinetrione derivatives and developed a structure-activity relationship (SAR). We then set about to using computational docking and pharmacophore mapping and 3D-QSAR modeling to investigate which of the two major STAT3 domains, i.e., SH2 and DBD, this novel class of inhibitors bind to. With this approach, we expanded the SAR and conducted pharmacophore mapping and 3D-QSAR studies using both a cis-conformational alignment and a trans-conformational alignment. In parallel, we applied structure-based modeling studies, including binding site mapping, docking and the molecular mechanics generalized born surface area (MM-GBSA)-based refinement analysis, to explore the binding pocket and poses of this class of compounds. Finally, surface plasmon resonance (SPR) studies were used to interrogate the suggestion by the computational studies that the novel pyrimidinetrione STAT3 inhibitors bind to the DBD STAT3, but not the SH2 domain.

Results

In vitro screening for selective STAT3 inhibitor lead structures

To develop novel drug leads, more than 300 in-house compounds were screened for their inhibitory effects on STAT3-DNA binding by electrophoretic mobility shift assay (EMSA) in vitro at 10 μM. Two hits, 1 and 2, which shared a common pyrimidinetrione core structure were reported in STAT3-DNA binding assay as previously reported [23,24]. Nuclear extracts containing active STAT3 prepared from v-Src-transformed mouse fibroblasts (NIH3T3/v-Src) were incubated for 30 min at room temperature with or without increasing concentrations of the hits, prior to incubation for 30 min with the radiolabeled hSIE probe, which binds to STAT3 and STAT1 and subjected to EMSA analysis [23,24]. Both compound 1 and 2 exhibited a dose-dependent inhibition of STAT3-DNA binding activity with IC_{50} values of 2.5 μM and 3.80 μM, respectively (Figure 1A). These values represent a dramatic improvement over those reported STAT3 inhibitors [18]. For selectivity, nuclear extracts containing active STAT1 and STAT3 prepared from the epidermal growth factor (EGF)-stimulated mouse fibroblasts over-

![Figure 1](https://example.com/figure1.png)

**Figure 1**: Selective effects of compounds 1 and 2 on the activities of STAT3. (A) Structure of compounds 1 and 2. Pyrimidinetrione lead structure is highlighted. (B) Nuclear extracts of equal total protein containing activated STAT1 and/or STAT3 were pre-incubated with or without compounds 1 or 2 for 30 min at room temperature prior to the incubation with the radiolabeled hSIE probe that binds Stat1 and Stat3, and subjected to EMSA analysis. (C) Effects of compounds 1 and 2 on pancreatic cancer cells (Panc-1) that harbor constitutively-active STAT3 and STAT3 knockout mouse embryonic fibroblasts (MEFs/STAT3−/−).
Table 1: Structure and STAT3-DNA inhibitory activities of pyrimidinetrione derivatives.

<table>
<thead>
<tr>
<th>No.</th>
<th>R&lt;sub&gt;1&lt;/sub&gt;</th>
<th>R&lt;sub&gt;2&lt;/sub&gt;</th>
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<th>pIC&lt;sub&gt;50&lt;/sub&gt;</th>
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*a pIC<sub>50</sub> = -log(IC<sub>50</sub>)

*Indicates external test set compounds.
model suggests that the NH group on the barbiturate ring is a favorable hydrogen bond donor (Figure 3A). The hydrophobic features at R2 are considered favorable (Figure 3B); furthermore, the model suggests that an electron-withdrawing group on the linker of the two aromatic rings on R1 group favors inhibitory activity (Figure 3C). Compounds designed were then constructed to match the pharmacophore-based 3D-QSAR model for activity prediction.

### Binding site mapping on STAT3

STAT3 dimerization via reciprocal phosphotyrosine-SH2 interaction between two monomers is considered as a crucial step in STAT3 function. A majority of reported STAT3 inhibitors were designed as STAT3 dimerization disruptors via computational virtual screening on the pY-peptide (APpYLKT) binding "hotspot" on SH2 domain, which consists of the pTyr705 binding pocket (Arg609, Lys591, Glu612 and Ser613), the Leu706 binding subsite and a hydrophobic side pocket (bordered by Ile597, Leu607, Thr622 and Ile634). Other possible inhibitor binding pockets have not been well studied. Thus, we carried out a binding site mapping study using the Sitemap program (Sitemap, Schrödinger, LLC, New York, NY) [33,34], which searched and scored likely binding sites on the STAT3 protein structure (PDB:1BG1 with DNA removed). A DNA binding region on STAT3 DNA binding domain (B) was identified

<table>
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<th>PLS Statistic</th>
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<td>SD</td>
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RMSE: Root Mean-Squared Error; SD: Standard Deviation of Regression; $r^2$: Non-Cross Validated R-squared; $q^2$: Cross Validated R-squared; F statistic: Overall Significance of the Model; Pearson-R: Pearson Correlation Coefficient

Table 2: PLS statistics of the best QSAR model, obtained with the cis-conformational alignment.

Figure 2: Pharmacophore-based QSAR model with the cis-conformation. (A) Alignment of compounds with the 5-point pharmacophore. Plots of QSAR model predicted activity ($pIC_{50}$) against actual STAT3 inhibitory activities ($pIC_{50}$) of training set (B) and test set (C) compounds.

(A) H-bond donor  
(B) Hydrophobic  
(C) Electron withdrawing

Figure 3: 3D-QSAR model visualized with different features. (A) H-bond donor; (B) Hydrophobic; and (C) Electron withdrawing property. Light blue cubes surrounding the pharmacophore features represent favorability feature and red cubes represent unfavorability.
as a putative binding site with the highest SiteScore value of 0.990 and the largest cavity (white dots), revealing it as the best druggable site on STAT3. This is interesting information, and could be exploited to advantage, considering that transcription factors are among the most difficult druggable targets [28,29]. In contrast, the most commonly targeted SH2 domain (Figure 4A) was assigned a lower SiteScore of 0.662, suggesting that it is an unsuitable druggable site, shallow and narrow. This is a significant observation supporting the recent attention to identifying direct STAT3 DBD binding inhibitors [19] that could change attitudes in attempts at drugging STAT3 [35].

Docking and MM-GBSA rescoring

To examine the binding site of these pyrimidinetriones on STAT3 in light of the above theoretical prediction, we applied docking studies [36,37] of all our tested 45 compounds (Table 1) to both the pY705 binding site and DNA binding site (PDB:1BG1 with DNA removed). Compound poses were then ranked by the Glide GScores. The best-docked compound on STAT3 DNA binding domain and SH2 domain has the GScore value of -5.530 and -4.797, respectively. The highest ranked docking pose of each compound was then subjected to MM-GBSA free energy calculation (Prime, Schrödinger, LLC, New York, NY) which evaluates binding free energies ($\Delta G_{\text{bind}}$) by minimization and MD simulation of protein-ligand complexes to address the errors generated by using rigid protein structure in regular docking procedures. Theoretically, potent inhibitors with high affinity ligand-protein binding should generate stable complexes, which have lower

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**Figure 4:** Binding site mapping and MM-GBSA refined docking studies on STAT3. Binding site contour maps of druggable sites on STAT3 (A) SH2 domain and (B) DNA binding domain. White filled circles indicate the size of the site; the red lattice represents the site surface. STAT3 protein surface is represented by electrostatic potential. Red indicates negative charge, and blue indicates positive charge. DNA is represented by the pink helical structure. Plots of STAT3 inhibition activity (pIC$_{50}$) versus MM-GBSA binding free energy of compounds docked in SH2 domain (C) or the DNA binding domain (D) 

**Table 1**

<table>
<thead>
<tr>
<th>Compound</th>
<th>pIC$_{50}$</th>
<th>MM-GBSA Binding Free Energy (kcal/mol)</th>
</tr>
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<tr>
<td>Comp1</td>
<td>6.0</td>
<td>-5.530</td>
</tr>
<tr>
<td>Comp2</td>
<td>5.5</td>
<td>-4.797</td>
</tr>
<tr>
<td>Comp3</td>
<td>5.0</td>
<td>-4.797</td>
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binding energies. In the MM-GBSA rescored binding energy-activity plot, we observed a good correlation ($r^2=0.50$, with the removal of one outlier) between the MM-GBSA binding free energy and the DNA binding inhibitory activity of the pyrimidinetrione derivatives in the cis-isomer configuration at the DNA binding domain (Figure 4D) but not at the SH2 binding domain (Figure 4C). No good correlation was shown with the trans-isomer configuration at either site (Suppl. Figure 2). The docking score and binding free energy of Compound 1 in both DNA binding domain and SH2 domain is listed on Suppl. Table 2. These data indicate that the pyrimidinetrione derivatives are likely to bind at the DNA binding domain of STAT3 instead of the supposed SH2 binding hotspot.

The topmost pose of compound 1 (IC$_{50}$=2.50 μM) with the binding free energy ($\Delta$Gbind) of -72.591 kcal/mol is shown in Figure 5. The pyrimidinetrione ring of the cis-isomer inserted into the cavity and engaged in hydrogen bonding with Leu430 and Val432 (Figure 5A). The inner phenyl ring of the R$_{1}$ group participated in π-π interactions with part of Arg423. The chlorine anchored the R$_{1}$ group to Asn420. Therefore, the molecule is locked in a fashion that the R$_{1}$ phenyl ring pointed toward the binding DNA (DNA structure was brought back into the model after the docking in order to indicate the site), suggesting a preclusion of binding. In addition, the protein structure underwent a large conformational change upon binding of compound 1 (Figure 5C), especially involving loop residues Arg382, Ser465 and Asn466, which interact directly with DNA [38]. This significant conformational change was not observed with the docking of the trans-isomers at the same site (Suppl. Figure 3). While R$_{1}$ and the pyrimidinetrione core facilitate binding, the R$_{2}$ substituent in the cis-conformation appears to sterically obstruct the DNA from binding to STAT3. To compare with the recently identified STAT3 DBD binding inhibitor inS3-54A18 [19], we constructed the molecule and docked it to STAT3 DNA binding domain (DBD). The top-ranking pose binds to the domain with a weaker binding free energy ($\Delta$Gbind) of -59.607 kcal/mol. The Val432 and Gly380 stabilized the carbonyl group and hydroxyl group of inS3-54A18 respectively through hydrogen bonding interactions. The chlorine The chlorine atom anchored the molecule to Asn472 (Figure 5C). Protein conformational change was not observed after inS3-54A18 docking (Suppl. Figure 3B) suggesting that molecule inS3-54A18 might have similar mechanism of inhibition of sterically obstructing the DNA from binding to STAT3; compound 1 has better binding at the STAT3 DBD.

Therefore, a two-step synthesis process described in Scheme 1B was designed to modify the R$_{1}$ substituent to increase the steric bulk in order to improve competitive inhibition of DNA binding to STAT3. Compounds with modifications of the R$_{2}$ group were also designed to expand the structural diversity keeping drug-likeness in mind (Scheme 1A).

*In vitro evidence that designed molecules interact with STAT3 and selectively disrupt STAT3 binding to DNA, but do not disrupt STAT3 binding to cognate pTyr peptide motif*

Compounds synthesized were purified and tested for inhibitory activity at the concentration of 10 μM via EMSA analysis. As we expected, compounds 63-67 with bulky and hydrophobic groups at R$_{1}$, had the best inhibitory effects against STAT3-DNA binding (Figure 6A). However, the preceding studies do not demonstrate a direct binding, or the binding site on the STAT3 protein. To provide a more definitive evidence of direct binding to STAT3, biophysical studies were performed. Recombinant human GST-tagged STAT3 protein (Abcam, Cambridge, UK) was immobilized on a Biacore CM5 sensor chip for surface plasmon resonance (SPR) analysis of the binding of compounds 63-67 in gradient concentration as analytes. Association and dissociation measurements were made and the binding affinities of the five compounds for STAT3 were determined. The results showed dose-dependent curves of the compounds upon their addition to the immobilized recombinant STAT3 (Figure 6B), indicative of the binding of compound to and dissociation from STAT3 protein. All the compounds exhibited good KD values except for compound 63 and 64, which showed poor solubility in the assay buffer. These data very strong evidence of the direct binding of the pyrimidinetrione STAT3

![Figure 5: Docking poses of compound 1 and inS3-54A18 at STAT3 DNA binding domain. (A) Amino acid shifts induced by compound 1 binding indicated by the green carbon colored amino acid structures. DNA is represented by the pink ribbon and sticks structure. (B) Protein conformational changes shown by overlaying the original structure (gray) and conformation resulting after the MM-GBSA refined docking (cyan). (C) Compound 1 (purple) and inS3-54A18 (orange) docked in STAT3 DNA binding site. Protein surface is presented with electrostatic potential property (red for negative charge and blue for positive charge).](image-url)
inhibitors to STAT3. These latter studies demonstrate that the designed STAT3 inhibitors 63-67 putatively interact with STAT3 and block STAT3 DNA binding. Given the computational modeling observation that pyrimidinetrione derivatives interact with the STAT3 at DNA binding domain, we surmise that these compounds block STAT3 DNA binding activity by directly disrupting DNA binding, unlike many STAT3 inhibitors that are thought to bind to the SH2 domain. To verify that pyrimidinetrione STAT3 inhibitors directly disrupt STAT3-DNA binding, we chose compound 67 with the best KD value to set up an SPR assay. The 5'-biotinylated hSIE probe was coated on a SA sensor chip. Recombinant STAT3 protein at 0.25 μM with or without 67 (20 μM), as well as 67 (20 μM) alone were incubated for 30 min at room temperature before injecting for SPR analysis as analytes. The data showed that STAT3 protein binds to 5'-biotinylated-hSIE probe with a response unit of 10 and this interaction was dramatically disrupted by adding compound 67 (Figure 7A). Therefore, as we expected from the model, compound 67 inhibits STAT3 binding to DNA by directly binding to STAT3 protein and interferes with DNA binding.

We then extended the studies to verify that 67 does not disrupt the pTyr:STAT3 SH2 domain interaction and subsequently inhibit STAT3:STAT3 dimerization. A similar SPR inhibition assay was designed using biotinylated STAT3-derived pTyr peptide (Biotin-PPYLKTK). The biotinylated STAT3-derived peptide was immobilized on the Biocore SA sensor chip as a ligand; 0.25 μM STAT3 protein with or without 20 μM 67 were incubated at room temperature for 30 min and then subjected to SPR analysis as analytes. No significant interruption of STAT3-pTyr peptide binding was observed in the presence of 20 μM 67 (Figure 7B), which indicated that the binding site of compound 67 on STAT3 was not on the pTyr binding site (SH2 domain) as we predicted in computational studies. This is a very significant result as we can now concentrate our attention on optimizing these new STAT3 inhibitor class using the 3D structures of STAT3 DNA binding domain. This is one of the rare occasions, where computational modeling has been used to elucidate a binding site that has been supported by experimental, direct protein binding studies.
Discussion and Conclusion

Although several STAT3 inhibitors have been hypothesized to bind in the STAT3 SH2 domain, the binding sites for small molecule inhibitors of STAT3 have not been identified. STAT3 dimerization via reciprocal phosphotyrosine-SH2 domain interaction between two monomers is considered as a crucial step in STAT3 function. Therefore, inhibiting STAT3 dimerization by displacing the pY-peptide (APpYLKT) binding to its pocket on STAT3 monomer is a rational strategy to inhibit STAT3 function. The binding hot spot on the STAT3 SH2 domain consists of three pockets, pTyr705 binding site (Arg609, Lys591, Glu612 and Ser613), Leu706 binding site and a side pocket (Ile 597, Leu607, Thr622 and Ile634). Using structure-based computational methods including virtual screening and fragment-based drug design, a series of STAT3...
dimerization inhibitors, such as STA-21 [39], S3I-201 [40], STX-0119 [41], celecoxib [42] and C188-9 [43], were developed by targeting the STAT3 SH2 domain. However, recent studies have indicated that nuclear import can also take place constitutively and independently of STAT3 tyrosine phosphorylation [20] and the binding of unphosphorylated STAT3 directly to DNA has also been observed [21]. This means that these above mentioned inhibitors could still miss inhibiting STAT3 function and be ineffective against STAT3 driven cancers. Therefore, the traditional strategy of inhibiting STAT3 dimerization may not be an efficient strategy to eliminate functions in tumor cells. Recently, a curcumin-based STAT3 inhibitor HO-3867 was suggested to bind directly to the STAT3 DNA binding domain [44] unlike other STAT3 inhibitors. Meanwhile, only few studies have been accomplished in other STAT3 inhibition strategies or binding site verification, which all the more emphasizes the challenge to STAT3 inhibitor design.

In our study, we discovered a set of novel pyrimidinetrione STAT3 inhibitors with higher inhibitory activity (low micro molar range) than most of the reported STAT3 inhibitors. Interestingly, they selectively inhibit DNA binding to STAT3 over STAT1. In vitro cell viability assays also showed that the compounds tend to inhibit proliferation in cancer cell lines that harbor constitutively active STAT3, relative to STAT3 knockout mouse embryonic fibroblast (MEF) cells. This indicates that the pyrimidinetrione STAT3 inhibitors have potential for selective STAT3 targeting. The pharmacophore-based 3D-QSAR model suggested that the substituent at the on R2 position is critical for STAT3 inhibitory activity. Our structure-based binding site mapping studies suggested the DNA binding domain contains the possible binding pockets for the compounds. From the MMGBSA-refined docking studies, we observed a significant correlation between binding energy at the STAT3-DNA binding domain and STAT3 inhibitory activity when the pyrimidinetrione compounds were docked at the DNA binding site. This energy-activity correlation was not seen with compounds docked to the SH2 domain, suggesting that the pyrimidinetrione STAT3 inhibitors bind at STAT3 DNA binding domain. The docking poses of the most active molecule also suggested that introduction of a hydrophobic or sterically bulky group at R2 could block DNA binding to STAT3. According to both the QSAR and docking models' observations, new inhibitors were synthesized and tested to confirm their ability to inhibit STAT3. As we predicted, compounds with the 4-((4-chlorophenyl)thio)phenyl substitution at R2 and bulky group at R1 gained the most inhibitory activity. Biophysics studies provided solid data for compounds directly binding to the STAT3 protein with good Kd values. We further developed an inhibitory assay using SPR analysis to detect the inhibitory effect of the best synthesized compound, 67, against STAT3 binding to DNA. Importantly, no significant interruption was observed when compound 67 was used to disrupt the pYyr peptide binding to STAT3. These data together show a strong support of our computational prediction that these novel STAT3 inhibitors disrupted STAT3-DNA binding by directly binding to DNA binding site, unlike many STAT3 inhibitors, which disrupt STAT3 dimerization. In view of the above note that STAT3 dimerization inhibitors may miss the interaction of STAT3 with DNA as even non-phosphorylated STAT3 has been shown to bind DNA and activate transcription. This new class of STAT3 inhibitors may serve as important drug leads or probes for the design and studies of STAT3 DNA-binding site inhibitors that should hold better promise for agent directed at interfering with STAT3 transcription activation.

Experimental Section

Forty-five commercially available pyrimidinetrione derivatives (Table 1) with diverse structural features were chosen by inspection and purchased from ChemBridge (San Diego, CA). Unless otherwise indicated most chemicals and reagents were purchased from Sigma-Aldrich (St. Louis, MI) to follow up on an initial screening of an in-house compound library.

Nuclear extract preparation and electrophoretic mobility shift assay (EMSA). Analyses were carried out as previously described [15,40]. Briefly, the 3P-labeled oligonucleotide probe was used as hSIE (high affinity sis-inducible element from the c-fos gene, m67 variant, 5'GCTTCATTTCCCGTAAATCCCTA) that binds STAT1 and STAT3 with high affinity. Nuclear extracts containing activated STAT3 or STAT1 protein were pre-incubated with compound for 30 min at room temperature prior to incubation with the radiolabeled probe for 30 min at 30°C before subjecting to EMSA analysis. Bands corresponding to DNA binding activities were scanned and quantified for each concentration of compound and plotted as percentage of control (vehicle) against concentration of compound, from which the IC50 values were derived.

Effects on STAT3 knockout mouse embryonic fibroblasts

Cell lines and reagents: The human pancreatic (Panc-1) cancer cells and the STAT3 null mouse embryonic fibroblasts (MEFs STAT3-/-) have all been previously reported [45]. These cells were grown in DMEM containing 10% heat-inactivated fetal bovine serum (FBS).

Cyquant cell proliferation assay: The CyQuant cell proliferation assay (Invitrogen Corp/Life Technologies Corp) was used to evaluate the biological activities of compounds, as previously reported [45] and following the manufacturer’s instructions. Relative cell viability of the treated cells was normalized to the DMSO-treated control cells.

Binding site mapping

The crystal structure of STAT3β homodimer bound to DNA (PDB:1BG1) was imported into Maestro interfaced in the Schrödinger suite. After protein preparation, DNA and water molecules were removed from the structure. Binding site mapping was carried out on the entire protein surface using the SiteMap program (version 2.7, Schrödinger, LLC, New York, NY). The calculation started with placing a 1 Å grid of possible site points around the protein. Grid points in good van der Waals contact with the receptor in solvent-exposed regions enclosed by the receptor served as site points. Then the sites were mapped onto another grid to generate five visualization maps, including hydrophilic, hydrophobic, H-bond donor or acceptor sites. Possible binding pockets were sorted using the SiteScore program, which is constructed and calibrated to 1 with the average score for 157 investigated sites. Therefore, a score greater than 1.0 suggests a very promising site, while a SiteScore below 0.8 indicates a non-drug-binding site [33,34].

Docking

All compounds were constructed in cis- and trans-conformations and underwent energy minimization and ligand preparation using LigPrep (version 2.8, Schrödinger, LLC, New York), which generated possible ionization states, stereoisomers at pH 7 ± 2. The protein preparation wizard in Maestro was used to prepare the STAT3 protein structure (1BG1) imported from Protein Data Bank (PDB) to render it ready for docking. DNA and water molecules were removed from the structure.

Receptor grid generation and docking studies were carried out with the GLIDE program (version 5.9, Schrödinger, LLC, New York, NY) [36,37]. A receptor grid on the SH2 domain, which can accommodate ligands with length up to 20 Å, was also generated by selecting residues Val637 and Ile 628, located in the center of the SH2 binding pocket, as
the centroid. Similarly, the docking grid at the DNA binding domain was created by picking amino acid residues Val342, Arg382, Gln415, Ser465, Asn466, which were within 3 Å distance from the DNA. Extra-precision (XP) mode of Glide docking was executed using a flexible ligand and a rigid receptor routine [46]. Compounds constructed in cis- and trans-conformations were docked to two binding sites identified by SiteMap. Ten docking poses were obtained from each MM-GBSA Rescoring. Compounds successfully docked were applied to a docking refinement process using the Molecular Mechanics/ Poisson-Generalized Born Surface Area (MM-GBSA) implemented in Prime MM-GBSA (version 3.2, Schrödinger, LLC, New York, NY). The analysis, which evaluates binding free energies by minimization and molecular dynamics (MD) simulation of the protein-ligand complex, were utilized to refine the conventional docking scores. The MM-GBSA binding free energy (ΔGbind) between a ligand and its receptor is calculated by the following equation.

\[ ΔG_{\text{bind}} = ΔH - TΔS - ΔEMM + ΔGsolv - TΔS \]

where ΔEMM is the MM energy difference between the ligand-receptor complex and the sum of the energies of free ligand and unliganded protein, ΔGsolv is the corresponding difference in the solvation energies and -TΔS is the corresponding difference in conformational entropy upon binding.

Pharmacophore perception

The 45 pyrimidinetrione derivatives and their observed activities were expressed as the negative logarithm of measured IC50 (pIC50) (Table 1) were used in this analysis. To facilitate the study of isomer influences, each compound was built in cis- and trans-conformations to generate two different pharmacophore models, respectively. A maximum of 1000 possible conformers (100 conformers per rotatable bond) were generated for each molecule using the MacroModel torsional sampling method (Mixed MCCM/IMOD), followed by 100 steps of minimization using MMFFs force field in a continuum solvent model. A set of pharmacophore feature sites, including hydrogen bond acceptor (A), hydrogen bond donor (D), hydrophobic group (H), negatively charged group (N), positively charged group (P) and aromatic ring (R), were assigned to each molecule. For both models, activity thresholds of 5.40 and 4.00 were set to identify four actives 2, 9, 10 and 28 and four inactive compounds 8, 14, 15, 33. A pharmacophore with five features common to all the four actives were identified using a tree based partitioning algorithm with a maximum tree depth of five. The pharmacophore box size, which governs the tolerance to match a pharmacophore, was set as 1 Å. The resulting common pharmacophore hypotheses were then ranked by survival score, which represents how well the active compounds superimpose when they are aligned on the features associated with that hypothesis.

Development of 3D-QSAR models

The top ranking common pharmacophore hypotheses was used to align molecules and generate atom-based 3D-QSAR models. The 45 pharmacophore aligned compounds with varying STAT3-DNA binding inhibitory activities were divided as training set and external test set to generate and validate the QSAR model, respectively. The ten test set molecules 2, 8, 12, 16, 18, 23, 36, 42, 44 and 45, were chosen in such a way that they represented the overall biological activity ranges, including the most active molecules, moderately active, and less active molecules. The rest of the molecules, 35 in total, were considered as a training set. The spread of activity (pIC50) in 3D-QSAR models covered approximately 2.0 logs range. In atom-based QSAR, a molecule is treated as a set of overlapping van der Waals spheres. The radii of these van der Waals spheres depend on the atom type in a ligand. Each atom of the ligand is classified into one of the six feature classes, comprising hydrogen bond donor, hydrophobic, negative ionic, positive ionic, electron withdrawing and miscellaneous. A rectangular grid is defined to encompass the space occupied by the aligned ligands. This grid divides the occupied space into consistently sized cubes for dimension of 1 Å, which are occupied by the atoms from the molecule. Therefore, each ligand can be represented by a set of binary-valued (0 or 1) to indicate the occupancy of the cube from different atoms. These occupancies of cubes and atom types are used as independent variables in generation of partial least squares (PLS) regression-based QSAR model [31]. All models were then validated by predicting the activity of the external test set molecules.

Surface plasmon resonance (spr) analysis

Biacore T200 (GE Healthcare, UK) was used for real-time binding interaction studies. For STAT3 and compounds binding analyses, a slow, high-level immobilization of recombinant human GST-tagged STAT3 (ab43618, Abcam, UK) protein was immobilized onto a CM5 series chip (GE Healthcare). The STAT3 protein was diluted to a concentration of 20 μg/mL in 10 mM Acetate buffer (pH=5.0). A 1:1 mixture of N-hydroxysuccinimide and N-ethyl-N-(dimethyaminopropyl) carbodiimide was used to activate 2 flow-cells of the CM5 chip. One flow-cell was used as a reference and thus immediately blocked upon activation by 1 M ethanolamine (pH=8.5). The sample flow-cell was injected with the diluted STAT3 at a flow rate of 10 μL/min. The STAT3 injection was stopped when the surface Plasmon resonance reached approximately 8000 RU. The analytes (compounds) were diluted in 1x PBS+1% DMSO buffer (filtered -0.22 um). The analytes were sequentially injected at a flow rate of 20 μL/min for 120s at 25°C, the dissociation time was set for 2 min. The concentrations were 6.25, 12.5, 25, 50, 100 μM for 63, 64, 3.125, 6.25, 12.5, 25, 50, 50 μM for 65–67. The equilibrium dissociation constant (KD) was obtained to evaluate the binding affinity by using the BIAEvaluation 2.0 software (GE Healthcare) supposing a 1:1 binding ratio. At least 3 independent experiments were performed.

For STAT3-DNA inhibition and STAT3-pTyr peptide inhibition assay, the biotin-hSIE probe (5'-biotin-AGCTTCATTTCCCGTAAATCCCTA, ThermoFisher, USA) or biotin-pTyr peptide (biotin-PpYLKTKP, Peptide 2.0, Chantilly, VA, USA) was immobilized onto a SA (streptavidin) chip (GE Healthcare). The biotin-hSIE (0.25 μM) or peptide-pTyr peptide (0.25 μM) was diluted in HBS-N and injected at a flow rate of 10 μL/min. The STAT3 flow-cell was used as a reference and thus immediately blocked upon injection of 20 ug/mL in 10 mM Acetate buffer (pH=5.0). A 1:1 mixture of biotin-hSIE protein was immobilized onto a CM5 series chip (GE Healthcare). The STAT3 protein was diluted to a concentration of 20 μg/mL in 10 mM Acetate buffer (pH=5.0). For STAT3 interaction studies. For STAT3 and compounds binding analyses, a slow, high-level immobilization of recombinant human GST-tagged STAT3 (ab43618, Abcam, UK) protein was immobilized onto a CM5 series chip (GE Healthcare). The STAT3 protein was diluted to a concentration of 20 μg/mL in 10 mM Acetate buffer (pH=5.0). A 1:1 mixture of N-hydroxysuccinimide and N-ethyl-N-(dimethyaminopropyl) carbodiimide was used to activate 2 flow-cells of the CM5 chip. One flow-cell was used as a reference and thus immediately blocked upon activation by 1 M ethanolamine (pH=8.5). The sample flow-cell was injected with the diluted STAT3 at a flow rate of 10 μL/min. The STAT3 injection was stopped when the surface Plasmon resonance reached approximately 8000 RU. The analytes (compounds) were diluted in 1x PBS+1% DMSO buffer (filtered -0.22 um). The analytes were sequentially injected at a flow rate of 20 μL/min for 120s at 25°C, the dissociation time was set for 2 min. The concentrations were 6.25, 12.5, 25, 50, 100 μM for 63, 64, 3.125, 6.25, 12.5, 25, 50 μM for 65–67. The equilibrium dissociation constant (KD) was obtained to evaluate the binding affinity by using the BIAEvaluation 2.0 software (GE Healthcare) supposing a 1:1 binding ratio. At least 3 independent experiments were performed.

Chemistry

Compounds were visualized by UV light on silica gel coated thin-layer chromatography plates. Flash column chromatography was performed on Fisher silica gel. 1D NMR spectra were recorded on Brucker 300 or 400 MHz NMR spectrometer, using dimethyl sulfoxide (DMSO) as solvent. Melting points of all products were determined using a Fisher-Johns melting apparatus. Mass spectra were obtained on a Brucker-HP 6000 spectrometer. Analytic high performance liquid chromatography (HPLC) was carried out using a SUPELCOSIL 5 µm C18 reverse phase column (250 x 4.6 mm) at ambient temperature on a Waters'
2695 HPLC system equipped with the 996 photodiode array detector. An isocratic method comprising 100% methanol (solvent B) was used. Five-min run time was set at a flow-rate of 1.0 mL/min. Area% purity was calculated at 254 nm.

4-((4-Chlorophenyl)thio)benzaldehyde (52)

10 mmol 4-chlorobenzanethiol was dissolved in DMF (5 mL) and equimolar potassium carbonates were added. The resulting reaction mixture was stirred at room temperature for 15 min then 4-bromobenzaldehyde (10 mmol) was added and stirred at room temperature for 4 h. Reaction was monitored using thin layer chromatography. The product was purified by column chromatography. White solid, yield (79%); M. P. 205-207°C; 1H NMR (400 MHz, DMSO-d6) δ 9.94 (s, 1H), 7.88-7.80 (m, 2H), 7.56 (d, J=0.7 Hz, 4H), 7.38-7.30 (m, 2H); MS (ESI) m/z 217.0 [M–H]-; HPLC: tR 2.529 min, purity 99.7%.

1-(4-Fluorophenyl)pyrimidine-2,4,6(1H,3H,5H)-trione (59)

White solid, yield 29%; M. P. 208-210°C; 1H NMR (400 MHz, DMSO-d6) δ 8.54 (s, 1H), 7.45-7.35 (m, 2H), 7.11-7.00 (m, 2H), 5.83 (s, 2H); MS (ESI) m/z 220.7 [M–H]-; HPLC: tR 2.562 min, purity 99.7%.

1-(3-Fluorophenyl)pyrimidine-2,4,6(1H,3H,5H)-trione (60)

Yellow solid, yield 33%; M. P. 188-189°C; 1H NMR (400 MHz, DMSO-d6) δ 11.61 (d, J=19.0 Hz, 1H), 7.56 (dq, J=15.7, 8.0 Hz, 1H), 7.40-7.25 (m, 1H), 7.17 (p, J=11.6, 10.7 Hz, 2H), 3.78 (d, J=18.4 Hz, 2H); MS (ESI) m/z 221.1 [M–H]-; HPLC: tR 2.629 min, purity 98.4%.

1-(2-(Trifluoromethyl)phenyl)pyrimidine-2,4,6(1H,3H,5H)-trione (61)

White solid, yield 48%; M. P. 174-176°C; 1H NMR (400 MHz, DMSO-d6) δ 11.73 (s, 1H), 7.86 (t, J=9.0 Hz, 2H), 7.69 (s, 1H), 7.56 (d, J=7.9 Hz, 1H), 3.88 (dd, J=166.5, 21.1 Hz, 2H); MS (ESI) m/z 271.1 [M–H]-; HPLC: tR 2.979 min, purity 98.0%.

General procedure for the synthesis of 1-substituted barbituric acids (53-62)

Urea (15 mmol) and diethyl malonate (15 mmol) were added to 250 ml ethanol with NaOEt (15 mmol). The reaction mixture was stirred and refluxed overnight then brought to room temperature. The solvent was partially evaporated and 100 mL cold water was added to the residue, acidified with 1M HCl and placed in ice-bath overnight and the resulting precipitate was filtered off and dried to afford the pure compound.

1-Phenylpyrimidine-2,4,6(1H,3H,5H)-trione (53)

Brown solid, yield 65%; M. P. 261-263°C; 1H NMR (400 MHz, DMSO-d6) δ 11.51 (s, 1H), 7.44 (dq, J=14.3, 7.3 Hz, 3H), 7.24 (d, J=7.5 Hz, 2H), 3.74 (s, 2H); MS (ESI) m/z 202.9 [M+Na]+; HPLC: tR 3.001 min, purity 98.0%.

1-[(1,1’-Biphenyl)-4-yl]pyrimidine-2,4,6(1H,3H,5H)-trione (54)

White solid, yield 28%; M. P. 220-222°C; 1H NMR (400 MHz, DMSO-d6) δ 11.54 (s, 1H), 7.83-7.65 (m, 4H), 7.51 (dd, J=8.4, 6.9 Hz, 2H), 7.46-7.38 (m, 1H), 7.37-7.28 (m, 2H), 3.76 (s, 2H); MS (ESI) m/z 279.6 [M–H]-; HPLC: tR 2.136 min, purity 99.7%.

1-(Naphthalen-2-yl)pyrimidine-2,4,6(1H,3H,5H)-trione (55)

White solid, yield 18%; M. P. 198-200°C; 1H NMR (400 MHz, DMSO-d6) δ 11.33 (s, 1H), 7.86 (m, 2H), 7.76 (m, 2H), 7.50 (m, 2H), 7.36 (dq, J=7.5, 7.5, 1.5 Hz, 1H), 3.12 (s, 2H); MS (ESI) m/z 252.8 [M+Na]+; HPLC: tR 2.677 min, purity 99.3%.

1-(o-Tolyl)pyrimidine-2,4,6(1H,3H,5H)-trione (56)

Yellow solid, yield 39%; M. P. 225-227°C; 1H NMR (400 MHz, DMSO-d6) δ 11.55 (s, 1H), 7.44-7.23 (m, 7H), 7.18 (d, J=7.6 Hz, 1H), 3.98-3.59 (m, 2H), 2.10 (s, 3H); MS (ESI) m/z 216.9 [M–H]-; HPLC: tR 2.520 min, purity 99.8%.

1-(m-Tolyl)pyrimidine-2,4,6(1H,3H,5H)-trione (57)

White solid, yield 40%; M. P. 210-212°C; 1H NMR (400 MHz, DMSO-d6) δ 11.49 (s, 1H), 7.35 (t, J=7.6 Hz, 1H), 7.23 (dt, J=7.3, 1.3 Hz, 1H), 7.07-6.99 (m, 3H), 3.74 (s, 2H), 2.34 (s, 3H); MS (ESI) m/z 217.0 [M–H]-; HPLC: tR 2.661 min, purity 99.9%.

1-(p-Tolyl)pyrimidine-2,4,6(1H,3H,5H)-trione (58)

White solid, yield 30%; M. P. 238-239°C; 1H NMR (400 MHz, DMSO-d6) δ 11.48 (s, 1H), 7.26 (d, J=8.0 Hz, 2H), 7.11 (d, J=8.0 Hz, 2H), 3.73 (s, 2H), 2.35 (s, 3H); MS (ESI) m/z 217.1 [M–H]-; HPLC: tR 2.529 min, purity 99.7%.
1-Phenyl-5-((4-(pyridin-2-ylxoxy)benzylidene)pyrimidine-2,4,6(1H,3H,5H)-trione (49)

White solid, yield 50%; M. P. 274–276°C; 1H NMR (400 MHz, DMSO-d6) δ 11.69 (d, J=63.4 Hz, 1H), 8.36 (d, J=27.1 Hz, 1H), 8.15 (dd, J=18.6, 4.9, 2.1, 0.8 Hz, 1H), 7.97 (dt, J=39.2, 2.0 Hz, 2H), 7.90–7.80 (m, 2H), 7.57–7.40 (m, 4H), 7.37–7.28 (m, 3H), 7.19–7.13 (m, 1H), 7.08 (ddt, J=21.5, 8.3, 0.8 Hz, 1H); MS (ESI) m/z 434.1 [M–H], 416.1 [M–MeOH–H]; HPLC: tR 2.121 min, purity 99.0%.

1-Phenyl-5-((4-(pyridin-3-ylxoxy)benzylidene)pyrimidine-2,4,6(1H,3H,5H)-trione (50)

Yellow solid, yield 48%; M. P. 253–255°C; 1H NMR (400 MHz, DMSO-d6) δ 11.68 (d, J=50.7 Hz, 1H), 8.65–8.49 (m, 2H), 8.39 (s, 1H), 8.33 (dd, J=6.6, 2.3 Hz, 1H), 8.28–8.21 (m, 1H), 7.60 (dd, J=12.9, 8.4, 4.8 Hz, 1H), 7.51–7.44 (m, 3H), 7.28 (d, J=8.2 Hz, 1H), 7.21–7.15 (m, 2H), 7.13–7.00 (m, 2H); MS (ESI) m/z 488.0 (m, 2H), 7.56 (d, J=2.0 Hz, 2H), 7.53–7.48 (m, 2H), 7.33 (ddt, J=7.9 Hz, 4.7 Hz, 2.1 Hz, 2H), 7.08 (ddt, J=21.5, 8.3, 0.8 Hz, 1H), 2.10 (s, 3H); MS (ESI) m/z 448.3 [M–3H], 479.1 [M–MeOH–H]; HPLC: tR 2.17 min, purity 99.9%.

5-(4-((4-Chlorophenyl)thio)benzylidene)-1-(p-tolyl)pyrimidine-2,4,6(1H,3H,5H)-trione (67)

Yellow solid, yield 67%; M. P. 279–281°C; 1H NMR (400 MHz, DMSO-d6) δ 11.69 (d, J=63.4 Hz, 1H), 8.36 (d, J=27.1 Hz, 1H), 8.15 (dd, J=18.6, 4.9, 2.1, 0.8 Hz, 1H), 7.97 (dt, J=39.2, 2.0 Hz, 1H), 7.91–7.80 (m, 2H), 7.57–7.39 (m, 4H), 7.37–7.27 (m, 2H), 7.20–7.13 (m, 1H), 7.08 (ddt, J=21.5, 8.3, 0.8 Hz, 1H), 2.10 (s, 3H); MS (ESI) m/z 448.3 [M–H], 479.1 [M–MeOH–H]; HPLC: tR 2.17 min, purity 99.9%.

1-Phenyl-5-((4-(4-chlorophenyl)thio)benzylidene)pyrimidine-2,4,6(1H,3H,5H)-trione (68)

Yellow solid, yield 71%; M. P. 246–247°C; 1H NMR (400 MHz, DMSO-d6) δ 11.72 (d, J=55.8 Hz, 1H), 8.29 (d, J=26.0 Hz, 1H), 8.11–7.98 (m, 2H), 7.56 (s, 3H), 7.52 (d, J=3.3 Hz, 2H), 7.28 (ddt, J=8.9, 2.5, 1.2 Hz, 1H), 7.26–7.21 (m, 2H), 7.21–7.15 (m, 2H); MS (ESI) m/z 451.9 [M–H], 482.9 [M–MeOH–H]; HPLC: tR 2.27 min, purity 99.9%.

5-(4-((4-Chlorophenyl)thio)(benzylidene)-1-(3-fluoro phenyl)pyrimidine-2,4,6(1H,3H,5H)-trione (69)

Yellow solid, yield 46%; M. P. 233–235°C; 1H NMR (400 MHz, DMSO-d6) δ 11.98 (s, 1H), 8.36 (d, J=33.9 Hz, 1H), 8.10 (dd, J=56.0, 8.6 Hz, 2H), 7.92–7.80 (m, 2H), 7.75–7.63 (m, 2H), 7.61–7.50 (m, 4H), 7.31–7.17 (m, 2H), 2.10 (s, 3H); MS (ESI) m/z 501.0 [M–H], 533.1 [M–MeOH–H]; HPLC: tR 1.93 min, purity 99.72%.

Analysis

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References

An Integrated Computational and Experimental Binding Study Identifies the DNA Binding Domain as the Putative Binding Site of Novel Pyrimidinetrione Signal Transducer and Activator of Transcription 3 (STAT3) Inhibitors.


