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Rapid Discovery and Structure—Activity Relationships of Pyrazolopyrimidines That Potently Suppress Breast Cancer Cell Growth via SRC Kinase Inhibition with Exceptional Selectivity over ABL Kinase

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ABSTRACT: Novel pyrazolopyrimidines displaying high potency and selectivity toward SRC family kinases have been developed by combining ligand-based design and phenotypic screening in an iterative manner. Compounds were derived from the promiscuous kinase inhibitor PP1 to search for analogs that could potentially target a broad spectrum of kinases involved in cancer. Phenotypic screening against MCF7 mammary adenocarcinoma cells generated target-agnostic structure—activity relationships that biased subsequent designs toward breast cancer treatment rather than to a particular target. This strategy led to the discovery of two potent antiproliferative leads with phenotypically distinct anticancer mode of actions. Kinase profiling and further optimization resulted in eCF506, the first small molecule with subnanomolar IC50 for SRC that requires 3 orders of magnitude greater concentration to inhibit ABL. eCF506 exhibits excellent water solubility, an optimal DMPK profile and oral bioavailability, halts SRC-associated neuromast migration in zebrafish embryos without inducing life-threatening heart defects, and inhibits SRC phosphorylation in tumor xenografts in mice.

INTRODUCTION

Modern drug discovery programs typically begin with a screening campaign (e.g., biochemical, virtual, or biophysical) for agonists, antagonists, or inhibitors of a nominated target associated with a particular disease.1−3 After hit identification, subsequent chemical optimization is fundamentally based upon “on-target” potency.1 The generation of high-affinity ligands (so-called lead compounds) is followed by chemical refinement into derivatives of superior potency, selectivity, and desirable pharmacokinetic properties.1,5 Selected drug candidates are then validated in vivo and, upon verification of safety and efficacy, progressed to human trials.5 While the merits of this well-defined process are undeniable, including several major breakthroughs in anticancer therapy,6 this strategy is also associated with declining productivity in the pharmaceutical industry and limited success to tackle the most aggressive cancers of unmet therapeutic need.7,8 High attrition rates at late stages of drug development underlines that cancer heterogeneity across patients and adaptive drug resistance mechanisms are major obstacles for the development of effective and long-lasting anticancer targeted therapies.9−12 These challenges have stimulated out-of-the-box thinking in pharmacotherapy research (e.g., targeted polypharmacology,10 antibody-drug conjugates,11 innovative prodrug approaches,14−17 etc.) and the re-examination of the core principles of drug discovery in complex diseases.18−20 The rise of modern phenotypic drug discovery19,20 together with the use of more clinically relevant disease models to guide early drug development20 are representative examples of the paradigm shift initiated in the field to trigger a positive inflection point.

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Protein kinases are integral components of signal transduction cascades. They govern a wide range of basic intracellular functions and coordinate cell-to-cell and extracellular matrix-to-cell communication to modulate cell and tissue physiology. Consequently, their malfunctioning is directly linked to progressive diseases including cancer and inflammation. The success in the clinic of several anticancer kinase inhibitors has validated a number of kinases as oncotargets, while the increasing understanding of cancer cell biology has demonstrated the essential role of different kinases in tumor suppressor pathways (antitargets).

The vast majority of kinase inhibitors bind to the kinase adenosine triphosphate (ATP) pocket. Since all kinases (>500) necessarily possess this relatively well-conserved catalytic site, there is a great potential for cross-reactivity. In fact, even though most kinase inhibitors are developed from single target hypotheses, they typically display broad selectivity profiles which, in some cases, have resulted in unanticipated clinical applications (e.g., sorafenib). Inhibitor promiscuity may also be advantageous for anticancer therapy when off-target activities assist to address bioactivity issues related to pathway redundancies, molecular heterogeneity, or resistance mechanisms. However, if these activities result in the inhibition of antioncogenic pathways or lead to severe side effects, drug promiscuity becomes a major drawback. Paradoxically, some kinases may behave as a target or an antitarget depending on the cancer context. The expression of the activated fusion oncoprotein BCR-ABL is a genetic abnormality associated with chronic myeloid leukemia (CML), and ABL inhibitors (imatinib, dasatinib) are clinically used in chronic phase CML treatment. Also, ABL family kinases are abnormally activated in various solid tumors, supporting their involvement in oncogenesis. However, ABL (ABL1) and ARG (ABL2) have been found to negatively modulate breast cancer progression in vivo, indicating that ABL inhibition could be counterproductive for breast cancer treatment (= antitarget). This example serves to delineate the complexity of cancer etiology and highlights the necessity of developing kinase inhibitors with tailor-made pharmacodynamic profiles for the effective targeting of each cancer subtype. Unfortunately, despite significant investments in the development of kinase inhibitors and the biomedical knowledge compiled over several decades, our still limited understanding of cancer biology prevents us from anticipating and optimally targeting the complex orchestrated actions that generate, maintain, and progress most neoplastic processes.

Acknowledging these limitations, many research groups including ours are frontloading the collection of robust empirical data to progress anticancer drug development programs away from classical black-and-white anticancer target hypotheses to more unbiased and evidence-led strategies for hit selection and lead generation. Following that principle, in this manuscript we show that cooperative ligand-based design and phenotypic screening, complemented with biochemical assays and the use of published data (literature, patents, etc.), can be effectively applied to accelerate the generation of preclinical drug candidates. Our strategy builds on three wide-ranging hypotheses: (i) targeting the kinase ATP pocket with compounds derived from promiscuous kinase inhibitors can enable “rationally-biased” serendipitous discoveries; (ii) early optimization of drug-likeness can be concurrently applied to explore pharmacodynamic diversity; and (iii) phenotypic screening of chemically related compounds in designated models of cancer can generate target-agnostic structure–bioactivity relationships and tailor ligand optimization to particular cancer types/subtypes. By means of this pragmatic approach to anticancer kinase inhibitor discovery, target deconvolution of identified hits and leads is largely simplified, thereby assisting the mechanistic elucidation of the molecular targets and antitargets involved in the observed phenotype.

In this work we describe how the implementation of such an approach led to the discovery of a kinase inhibitor with potent activity against breast cancer cells and a unique selectivity profile: the first small molecule able to inhibit SRC at subnanomolar levels with a 1000-fold selectivity over ABL.

## RESULTS AND DISCUSSION

### Design, Synthesis, and Screening of Compounds 7a–I

PP1 is a promiscuous inhibitor that indiscriminately targets protein tyrosine kinases, many of which are involved in oncogenesis such as the SRC family kinases (SKF), RET, KIT, and ABL. Moreover, related derivatives developed thereafter have shown strong inhibition of a variety of kinases with relevance in cancer including IGF-1R, EGFR, BTK, VEGFR, PDGFR, P13K, and mTOR. According to the crystal structure of PP1 with HCK and RET kinases, this small molecule is an archetypical type I kinase inhibitor, with its N5 and 4-NH2 groups forming multiple H-bonds with the hinge region of the kinase ATP site (Figure 1). The C3 p-tolyl group is positioned toward a hydrophobic region well-conserved across tyrosine kinases, thus being responsible for the partial selectivity of PP1 over other kinase families. Although PP1’s potent inhibition of disease-associated kinases make it a valuable biological tool, its use is limited by low solubility in water and poor selectivity, major limiting factors for the clinical translation of many drug candidates.

We envisaged that the substitution of PP1’s tert-butyl group at the N1 position with flexible water-solubilizing groups could be used to improve drug-like properties and, at the same time, explore the accessible sugar/phosphate regions occupied by the natural ligand ATP in the search for novel binding affinity profiles. As shown in Figure 1, compounds were designed to display a cyclic tertiary amine connected to the 5 and 4-NH2 groups forming multiple H-bonds with the hinge region of the kinase ATP site (Figure 1). The C3 p-tolyl group is positioned toward a hydrophobic region well-conserved across tyrosine kinases, thus being responsible for the partial selectivity of PP1 over other kinase families. Although PP1’s potent inhibition of disease-associated kinases make it a valuable biological tool, its use is limited by low solubility in water and poor selectivity, major limiting factors for the clinical translation of many drug candidates.

![Figure 1](image-url)

**Figure 1.** Upper panel: ATP and PP1 (neutral forms). Lower panel: General structure of the novel compounds and retrosynthetic analysis.
the pyrazolopyrimidine ring through an ethylene linker. Following the route described in Scheme 1 (see full details in the Supporting Information), a highly focused 12-member library was generated by coupling a selection of cyclic secondary amines to an aldehyde-modified derivative 43 of PP1 (6) via reductive amination.

The 12-member library was then tested against human mammary adenocarcinoma MCF7 cells, using the inhibition of cell growth as the primary output of the screening. Live-cell high-content image-based phenotypic assays were carried out to kinetically profile the antiproliferative response of MCF7 breast cancer cells following treatment with a single dose (100 μM) of compounds 7a–l for 5 d. Experiments were performed in triplicate, using cells treated with PP1 (100 μM) or DMSO as a positive and negative control, respectively. Cell proliferation was monitored by time-lapse imaging using an IncuCyte ZOOM microscope and analyzed by its software, enabling determination of cell density (% confluence) over time after compound addition. As shown in Figure 2a, compounds 7d and 7i–l (colored lines) led to a strong inhibition of cell proliferation, maintaining confluence levels equal or below those imaged at time zero (point of compound addition). Interestingly, all these hits presented at least two tertiary amines in their N1 motifs, providing the first insights into structure/antiproliferative activity relationships.

Half-maximal effective concentration (EC₅₀) values were then calculated for the five hits and PP1 in MCF7 cells using an 8-point half-log dose–response study (0.03 to 100 μM). Based on the MCF7 growth kinetics data provided by the IncuCyte assay, cell viability was determined at day 5 using the PrestoBlue reagent. Figure 2b shows that derivatives 7j and 7k were the most potent among the novel compounds, with EC₅₀ values of 12 μM. Due to its potency and lower molecular weight, the dimethylamino-containing piperidinyl group of compound 7j was chosen as the N1 motif for the preparation of a second library of pyrazolopyrimidines.

**Design, Synthesis, and Screening of Compounds 8a–f and 9a–f.** Several investigations have reported that substitutions of the p-tolyl group at the C3 position of PP1 by different aryl moieties (even closely related ones) have a major impact on protein–ligand binding. medicinal chemistry at that position has generated inhibitors for a variety of kinases, including receptor and nonreceptor tyrosine kinases (e.g., SRC, ABL, RET, EGFR, BTK, PDGFRs, VEGFRs, KIT)34–38,40 and nontyrosine kinases (e.g., PI3Ks, mTOR).39,44 To exploit this feature as an opportunity to expand the prospective pharmacodynamic scope of 7j, a selection of five arylboronic acids and phenylacetylene were employed to functionalize the C3 position of the heterocyclic ring by palladium-catalyzed cross-coupling chemistry to synthesize derivatives 9a–f.
Together with the piperidine-containing compounds 9a−f, the corresponding acetal intermediates 8a−f were selected for testing against breast cancer cells to increase the chemical diversity of the screening.

Antiproliferative properties of compounds 8a−f and 9a−f were evaluated in MCF7 cells (dose–response studies), using compound 5 (acetal intermediate of library 1), 7j (most potent derivative of library 1), and PP1 as controls. As shown in Figure 3, compounds 8d and 9d (both having a 7-azaindol-5-yl group at C3) exhibited superior antiproliferative properties (EC$_{50}$ < 2 μM). Remarkably, time-lapse imaging of compounds 8d and 9d (1−100 μM) revealed patently distinct antiproliferative mode of actions (Figure 3b and Movies S1 and S2). Compound 9d—containing the piperidinyl moiety of 7j—induced cell death, while its acetal precursor 8d halted cell division.

To identify the target/s that could be responsible of the phenotype induced by hits 8d and 9d in MCF7 cells and thereby shed some light over further optimization campaigns, inhibition activities were tested for PP1, 7i−k, 8d, and 9d against a selection of kinases involved in human cancer. Proteins were chosen in accordance with the profile of related pyrazolopyrimidines. Kinase profiling (Reaction Biology Corp., USA) was performed by measuring $^{33}$P incorporation on the substrate (poly [Glu, Tyr] 4:1) relative to DMSO. Dose–response curves and calculated half-maximal inhibitory concentration (IC$_{50}$) values are shown in Figure S1 and Table 1, respectively. Derivatives 7i−k and 9d strongly inhibited SFK members, whereas acetal-functionalized compound 8d preferentially inhibited mTOR (>28-fold more potent than PP1). The most potent inhibitor was compound 9d, which displayed similar potency to PP1 against SRC but higher selectivity over RET, PDGFRα, and KIT. Interestingly, according to the inhibition profile of compounds 7i−k (only varying on the spacer between the dimethylamino group and the piperidine), potency and selectivity for SRC were enhanced by the proximity of the dimethylenamino to the ring (IC$_{50}$(ABL)/IC$_{50}$(SRC) for 7i = 50).

Using the information provided by the phenotypic screening and the kinase inhibition profiling of leads 8d and 9d, two independent optimization campaigns were implemented toward the generation of anticancer drug candidates inhibiting mTOR and SFK, respectively. Optimization of lead 8d into the highly selective mTOR inhibitor eCF309 has been reported elsewhere. Herein is described the generation and structure–activity relationships (SAR) of pyrazolopyrimidine derivatives with high potency and selectivity for SFK.

**Design, Synthesis, and Phenotypic Screening of Compounds 11a,b.** The data collected from the kinase profiling study indicated that the 2-[4-(dimethylamino)-1-piperidyl]ethyl group at the N1 position was optimal to generate SFK inhibitors of high selectivity, while the superior potency of 9d against SRC suggested that on-target potency could be further enhanced by optimization of the substituent at the C3 position. A literature survey on SRC inhibitors was thus performed to assist in the design of derivatives with improved SRC binding. We focused our attention on the C3 substituent of PP20, a potent dual SRC/ABL inhibitor developed by Apsel et al. (Figure 4a). Novel derivatives were designed by substituting the 7-azaindol-5-yl group at C3 of 9d with the functionalized phenyl ring found in PP20, whereas the piperidines used for 7i and 7j were introduced at the N1 position.
position. Acetal deprotection of compound \(4\) in TFA/water (1:1) followed by reductive amination with either 4-(dimethylamino) or 4-(dimethylaminomethyl)piperidine generated iodo intermediates \(10a, b\), respectively, which afforded derivatives \(11a\) (also known as eCF506) and \(11b\) (Figure 4a) by Suzuki cross-coupling with 4-(N-Boc-amino)-3-methoxyphenylboronic acid (see synthesis in the Supporting Information).

Antiproliferative properties of compounds \(11a, b\) were then tested in cells using PP20 and dasatinib (a clinically approved dual SRC/ABL inhibitor)\(^{22}\) as positive controls. Along with MCF7 cells, breast adenocarcinoma MDA-MB-231 cells were tested as a model of triple-negative breast cancer known to be particularly sensitive to SRC inhibitors.\(^{46}\) As shown in Figure 4b, compound \(11a\) induced a very potent antiproliferative effect in both MCF7 and MDA-MB-231 cells, significantly superior to the activity displayed by derivatives \(9d, 11b,\) and PP20. Notably, \(11a\) also outperformed the gold-standard SRC inhibitor dasatinib.

Compound \(11a\) was then selected for further screening. Inhibition of SRC kinase activity in cells was studied by Western blot. Upon activation, SRC undergoes autophosphorylation of the Y416 residue within its kinase domain.\(^ {47}\) Study of phosphorylated SRCY416 levels can then serve both as evidence for the presence of active SRC and to evaluate direct inhibition of SRC kinase activity. Phosphorylation of the focal adhesion kinase (FAK), a downstream substrate of SRC with high relevance in cancer progression,\(^ {48}\) was also studied in both MDA-MB-231 (Figure 4c) and MCF7 (Figure S2) cells. Dasatinib was used as a positive control. Cell lysates were prepared following 24 h of serum starvation and 1.5 h exposure to each inhibitor and serum stimulation for 1 h. Analyses demonstrated that \(11a\) inhibits phosphorylation of SRC and FAK at low nanomolar levels, with complete inhibition observed at 100 nM. Dasatinib

Table 1. IC\(_{50}\) Values (in nM) Calculated for PP1, 7i–k, 8d, 9d, 11a, and Dasatinib in a Selection of Recombinant Kinases

<table>
<thead>
<tr>
<th>kinase \ hit</th>
<th>PP1</th>
<th>7i</th>
<th>7j</th>
<th>7k</th>
<th>8d</th>
<th>9d</th>
<th>11a</th>
<th>dasatinib</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABL</td>
<td>147</td>
<td>6323</td>
<td>7525</td>
<td>7249</td>
<td>1207</td>
<td>116</td>
<td>479</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>FYN(^ {a})</td>
<td>27</td>
<td>311</td>
<td>913</td>
<td>1964</td>
<td>1226</td>
<td>38</td>
<td>2.1</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>KIT</td>
<td>1318</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>39</td>
</tr>
<tr>
<td>mTOR</td>
<td>9318</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>328</td>
<td>8565</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
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<tr>
<td>PDGF(\alpha)</td>
<td>657</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
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<tr>
<td>RET</td>
<td>11</td>
<td>6584</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>&gt;10(^ 4)</td>
<td>433</td>
</tr>
<tr>
<td>SRC(^ {a})</td>
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<td>303</td>
<td>1040</td>
<td>2453</td>
<td>27</td>
<td>&lt;0.5</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>YES(^ {a})</td>
<td>36</td>
<td>71</td>
<td>344</td>
<td>696</td>
<td>566</td>
<td>12</td>
<td>&lt;0.5</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>IC(<em>{50})(ABL) /IC(</em>{50})(SRC)</td>
<td>8</td>
<td>50</td>
<td>25</td>
<td>7</td>
<td>0.5</td>
<td>4</td>
<td>&gt;950</td>
<td>1</td>
</tr>
</tbody>
</table>

\(^{a}\)Member of SFK.

Figure 4. (a) Structures of PP20 and compounds \(11a\) and \(11b\). (b) EC\(_{50}\) values calculated after treating MCF7 and MDA-MB-231 cells with compounds \(11a, 11b, PP20,\) and dasatinib (dose range: 1–10,000 nM) for 5 d. Cell viability was determined using the PrestoBlue reagent. Error bars: ± SD from \(n = 3\). (c) Western blot analysis of SRC and FAK activity in MDA-MB-231 cells treated with \(11a\) and dasatinib for 1.5 h. (d) Scratch-wound migration assay. MDA-MB-231 cells were treated with \(11a\) or dasatinib (10 nM), and cell migration compared with untreated cell control (DMSO, 0.1% v/v) at 6, 12, and 24 h. Cells were imaged and analyzed using an IncuCyte-ZOOM microscope with integrated scratch-wound migration software module. Error bars: ± SD from \(n = 3\); \(p < 0.001, ***; p < 0.01, **.\) P value calculated from t-test. (e) Snapshots of the scratch-wound area of MDA-MB-231 cells at 0 and 24 h. Yellow lines highlight the gap formed by the scratch. Blue lines indicate the advance of cells into the wound after 24 h incubation in the absence and presence of \(11a\) (10 nM).
activity was close to that of 11a, although it also induced a dose-dependent increment of total SRC (Figure 4c).

Having demonstrated that 11a targets SRC in cells, we investigated FAK activity and cell viability in cells that lack expression of SRC. SYF murine embryonic fibroblasts (SRC−/−, YES−/−, FYN−/−) were treated with either 11a or dasatinib and levels of total/phospho-FAK analyzed by protein immunoblot. As expected (Figure S3), FAK activity was found to be unaffected by drug treatments, indicating that the inhibition of phospho-FAK in the breast cancer cell lines was a consequence of SRC inhibition. Compounds 11a and 11b displayed significantly less antiproliferative activity in SYF cells than dasatinib, suggesting increased selectivity against SRC family kinases (Figure S3).

An automated 96-well scratch-wound cell migration assay (analyzed in real-time by the IncuCyte ZOOM system) was then set up to determine whether compound 11a could halt migration of MDA-MB-231 cells, as would be expected for a SRC inhibitor.49 Cells treated with 11a were monitored for 24 h at a dose range (1−10,000 nM) and compared with untreated cells (0.1% v/v DMSO) and dasatinib treatment. Compound 11a significantly reduced cell motility at 10 nM as early as 6 h into the study, with equivalent efficacy to dasatinib (Figure 4d,e). 11a treatments at higher concentrations induced greater inhibition of cell migration (Figure S4), although cell viability was also significantly reduced. Image-based measurement of caspase 3/7 activity in MCF7 breast cancer cells demonstrated significant levels of apoptotic cell death after 5 d treatment with 11a at concentrations ≥100 nM (Figure S4), in accordance with the EC50 values observed for this cell line.

**Kinome Profiling of SRC Inhibitor 11a.** EC50 values were determined for 11a and dasatinib against the panel of proteins in MCF7 and MDA-MB-231 cell lines (Figure 5). These values were compared with the IC50 values against SRC and ABL kinase activities (as IC50) and were used to generate a profile of the compounds' selectivity and potency. The results showed that 11a exhibited a higher degree of selectivity for SRC over ABL, as indicated by the lower EC50 values for SRC inhibition compared to those for ABL (Figure 5).

**Figure 5.** Compounds 12a−x and their biological activities. Structural motifs modified on 11a for the generation of compounds 12a−x are highlighted as colored circles. EC50 values against MDA-MB-231 breast cancer cells are shown below each structure. SRC and ABL inhibition values (as IC50) are reported for selected compounds.
Figure 6. (a–d) Predicted binding modes of PP20 and 11a in SRC and ABL kinases. (a) PP20 in SRC, (b) PP20 in ABL, (c) 11a in SRC, and (d) 11a in ABL. The P-loop, αC helix, and DFG motif are in red, orange, and magenta, respectively. PP20 and 11a are represented by sticks, where carbon atoms are in black, nitrogen atoms are in blue, and oxygen atoms are in red. (e) 11a in SRC active site compared to adenosine monophosphate (AMP) (from PDB 3DQX). AMP is represented by sticks, where carbon atoms are in green, nitrogen atoms are in blue, oxygen atoms are in red, and the phosphorus atom in orange. Predicted binding mode indicates that the dimethylamino group at the piperidine ring is optimally placed to interact with the carboxylate group of Asp404 (DFG motif, in pink).

recombinant kinases used in previous screenings (see IC_{50} values in Table 1 and dose–response curves in Figure S5). As expected, dasatinib induced potent inhibition of nonreceptor tyrosine kinases, such as ABL and SRC, but also high potency against receptor tyrosine kinases (e.g., PDGFRα, KIT). In contrast, 11a exclusively inhibited SFK, with subnanomolar IC_{50} values against SRC and YES (activities for the rest of the SFK members are shown in Table S1). It is important to highlight that 11a displayed a vast difference in activity (>950-fold difference) between ABL and its primary target SRC. While small molecule inhibitors with reverse binding capabilities have been developed (e.g., imatinib strongly targets ABL without inhibiting SRC), this is, to the best of our knowledge, the first case of a small molecule with subnanomolar IC_{50} for SRC that requires a concentration 3 orders of magnitude greater to reach the same level of inhibition in ABL. Such properties have only been previously reported in peptide-based bisubstrate inhibitors of SRC. 

Design and Synthesis of Compounds 12a–x and SAR Analysis. To shed light over the structural features responsible for the high antiproliferative properties and unique selectivity profile of 11a, a library of 24 closely related analogs was developed (see syntheses in the Supporting Information) by introducing small changes on key functional groups of 11a (see Figure S5). The activity of compounds 12a–x was then tested in cells. MDA-MB-231 was the cell line chosen for the screening due to its superior sensitivity to 11a treatment. Cell viability assays were performed as previously described, and EC_{50} values calculated for each compound. IC_{50} values were subsequently determined against SRC and ABL for those compounds exhibiting high antiproliferative activity (12d, 12e, 12m, 12w, and 12x) and for selected compounds with low activity (12a, 12i, 12j, 12t, and 12v).

Analysis of the properties of compounds 12a–e (Figure 5) indicates that the presence of a tertiary amino group at the position 4 of the pyperidinyl ring is essential for the activity. Substitution of the methyl groups by larger aliphatic rings was tolerated, however, with compounds 12d and 12e being among the most potent SRC inhibitors of the series. Removal of the pyrimidine ring (see compound 12f) resulted in >3000-fold decrease in antiproliferative activity. Screening of compounds 12g–x evidenced the limited structural variations permitted at the top of the molecule to achieve high bioactivity. Most modifications of the Boc and the methoxy groups of the phenyl moiety at the C3 position led to a significant reduction in bioactivity. Even minor chemical modifications such as the substitution of the carbamate group by urea (12i) or amide (12j), or the change of the methoxy by OH (12s), resulted in >200-fold decrease in both antiproliferative properties and SRC inhibitory activity. Remarkably, introduction of a benzylamino group instead of the Boc was tolerated well, as observed from the bioactivity exhibited by compound 12m. However, introduction of endocyclic nitrogen atoms in the ring (see 12n and 12o) dramatically reduced compounds’ activity, particularly in the ortho position. Interestingly, while the methoxy group is required for maintaining high levels of activity when the Boc group is present in the structure (see 12s–u), substitution of the Boc group by an ester tolerated the elimination of the methoxy group (see 12w and 12x).

Biochemical and in Silico Studies of 11a Binding to the SRC Kinase. Most kinase inhibitors, especially adenine analogues, bind to the catalytic domain of the enzyme in its active conformation, thus competing with the natural substrate ATP. To determine whether this was the mode of binding of 11a to its primary target SRC, competitive inhibition studies with ATP were performed (Reaction Biology Corp., USA). Experiments were designed to obtain K_{m} values for ATP in the absence and presence of different concentrations of inhibitor 11a. The reactions were monitored every 10 min to obtain progress curves with time course (Figure S6). These were found to be linear regardless of the compound’s concentration, suggesting that SRC inhibition by 11a is not time-dependent (= reversible). The slopes (µM/min) were represented against ATP concentration for a Michaelis–Menten plot and a subsequent Lineweaver–Burk plot (double-reciprocal plot), using GraphPad Prism software.
Compounds were added to 2 dpf zebrafish (500 μM) and imaged at 72 hpf. (a) Imaging analysis of the distance between the last neuromast and the tip of the tail (n = 3) under treatment with DMSO (negative control) or 11a (500 μM). P value calculated from t-test. (c) Study of zebrafish heart development under short treatment with 11a (500 μM) and dasatinib (10 μM). Compounds were added to 2 dpf zebrafish embryos and incubated for 4 h (n = 3). Subsequently, fresh media was added, and the fish imaged after 48 h incubation.

To gain insight on the selective binding affinity of 11a for SRC over ABL, in silico docking studies were performed with both the dual ABL/SRC inhibitor PP20 and the SRC-selective inhibitor 11a. Docking of PP20 into SRC and ABL produced a plausible predicted binding mode in both cases (Figure 6a,b), with the adenine-mimicking pyrazolopyrimidine making H-bonds to the hinge region of the proteins. The tert-butyl group of the carbamate occupies a hydrophobic pocket formed by several lipophilic residues in SRC and ABL, including the Phe of their aspartate-phenylalanine-glycine (DFG) motif. Free energy of binding was predicted by Autodock to be approximately the same for both proteins (−8.5 and −8.9 kcal/mol, respectively). Docking of 11a into SRC also produced a plausible predicted binding mode (Figure 6c) and a higher predicted affinity than PP20 (calculated free energy of binding of −10.6 kcal/mol). In SRC, the dimethylamino group of the ethylpiperidinyl moiety on N1 interacts with Asp404, one of the protein’s phosphate binding residues (see Figure 6c,e). In contrast, the large N1 group of 11a (compared to the isopropyl on PP20) clashes with Tyr253 in the catalytic site of ABL. This resulted in Autodock being unable to find a position in the ATP binding site of ABL where 11a could mimic the interactions made by the adenine moiety of ATP (Figure 6d). This study indicates that the flexible polynamine linker on N1 is responsible for the low affinity of compound 11a for ABL.

Phenotypic Screening of SRC Inhibitor 11a in Zebrafish. Developing zebrafish provides a rapid phenotypic assay to simultaneously test safety and efficacy of novel compounds in a living vertebrate. Small molecule phenotypic-based screens in zebrafish have recently implicated SRC kinase in the migration of the posterior lateral line primordium, a cohesive cluster of cells that migrates horizontally under the skin along the myoseptum to the end of the tail, periodically depositing neuromasts. To determine the effects of 11a on cell migration in vivo, we treated Tg(bm3:cmGFP) transgenic zebrafish that express green fluorescent protein (GFP) in the mechanosensory hair cells of the lateral line (which form part of the neuromasts) with 11a for 2 d and measured the distance of the last neuromast to the tip of the tail (marked by the end of the notochord and the presence of black melanocytes, Figure 7a, in red). 11a significantly reduced neuromast migration (>100 μm in average) with minimal effect on the development of the embryos (Figure 7a–c). In contrast, dasatinib treatment at >10 μM resulted in severe cardiotoxicity and death of most embryos. At concentrations that were compatible with embryo survival (1–10 μM), dasatinib did not inhibit the migration of neuromasts, whereas it did still induce a patent cardiotoxic phenotype (note heart enlargement in Figure 7c). Further safety studies (see Figure S8) showed that dual ABL/SRC inhibitor PP20 also induces severe cardiotoxicity in zebrafish even after short treatment. These results, which correlate with the essential role of ABL in heart development and healing, suggests that the selectivity of 11a over ABL might be advantageous for therapy when ABL inhibition is not required.

Physicochemical and Drug Metabolism and Pharmacokinetics (DMPK) Profiling of SRC Inhibitor 11a in Vitro and in Vivo. Compound 11a displays very high solubility in water (>100 mg/mL, pH = 7.4), a significant advantage over dasatinib, whose solubility in water ranges from moderate (18 mg/mL at pH = 2.6) to extremely low (<0.001 mg/mL at pH = 7). Notably, analysis of cLogP with the OSIRIS Property Explorer provided an estimated value of 2.2 for 11a, whereas experimental log D at pH = 7.4 was determined to be −0.04. Inhibition of hERG channel and cytochrome P450 (CYP) enzymes was determined for 11a to assess off-target liabilities associated with cardiotoxicity (QT interval prolongation) and potential drug–drug metabolic interactions. Results demonstrated weak hERG inhibition (50.1% inhibition at 25 μM) and marginal inhibition of CYP enzymes at 10 μM (Figure S9).

A plasma protein binding assay was performed with compound 11a to test to what extent the compound can bind to proteins in the blood, an important factor to predict free drug levels in the body. Warfarin (positive control, 99–100% bound) and 11a were incubated with human or rat plasma in duplicates at 37 °C for 24 h and the free (unbound) fraction of compound determined by LC-MS/MS. Percentages...
of unbound 11a were 9.4 and 19.1% in rat and human plasma, respectively.

In vitro metabolic stability of 11a was systematically assessed in human liver microsomes (HLM), blood plasma, and primary hepatocytes (Cyprotex, U.K.). LC-MS/MS analysis of 11a showed excellent compound stability (94%) after 30 min incubation with HLM at 37 °C. 11a was subsequently incubated (1 μM, 2 h, 37 °C) with plasma from either human, mouse, or rat and analyzed by LC-MS/MS at different time points. Regardless of the species, analyses showed full plasma stability (Figure S10). Lastly, compound 11a (3 μM) was incubated with mouse and human primary hepatocytes for 60 min and samples analyzed by LC-MS/MS at various time points. Verapamil and umbelliferone were used as control compounds. In agreement with the HLM study, compound 11a showed high stability in human hepatocytes with a half-life of 751 min (Figure S10). As expected, faster clearance was observed in mice hepatocytes (7-fold decrease in half-life).

Encouraged by the in vitro DMPK profile of 11a, an in vivo study was performed in mice to determine oral bioavailability and half-life (Cyprotex, U.K.). Female CD1 mice were given a single dose of 10 mg/kg (0.25 mg/25 g mouse) of compound 11a orally (as a solution in nanopure water) or via IV injection (as a saline solution). Blood/plasma samples were taken at different time points (final sample taken after 8 h), and compound levels measured by LC-MS/MS (Figure S11). The calculated half-life value for 11a was 2.9 h, substantially higher than that reported for dasatinib in mice (0.9 h).58 Compound 11a showed moderate oral bioavailability (25.3%), although it is important to note that no excipients (just pure water) were used for its oral formulation.

**In Vivo SRC Inhibition Study in a Tumor Xenograft Model.** The presence of active (phosphorylated) SRC in human colorectal cancer HCT116 cells and its inhibition under 11a treatment was verified by Western blot (Figure S12). Subsequently, an in vivo PD study was performed in a xenograft model of HCT116 cells in mice.59 HCT116 cells were injected subcutaneously, and tumors were allowed to grow up to 3–4 mm in diameter. Subsequently, mice were dosed daily for 3 d with 11a (50 mg/kg, in nanopure water) or vehicle (nanopure water) by oral gavage and culled 3 h after the last dose (n = 4). Tumors were excised, fixed, and sections labeled for phospho-SRCY416 and stained with hematoxylin. As shown in Figure 8, microscopy analysis demonstrated significant reduction of phospho-SRCY416 in the xenograft sections from mice treated with 11a relative to the untreated animal controls.

**CONCLUSIONS**

To move away from orthodox target-centric strategies, a pragmatic approach for rapid discovery of novel high-quality anticancer kinase inhibitors was implemented through the generation and phenotypic screening of analogues of PP1, a pyrazolopyrimidine known to inhibit multiple oncogenic pathways.12–34 Derivatives of this promiscuous kinase inhibitor were designed to improve physicochemical properties and explore favorable pharmacological features through small chemical modifications. To accelerate the advance from hits to leads to drug candidates, compounds were screened by phenotypic assays in the search for derivatives with potent antiproliferative properties. Chemical design was biased toward human breast cancer treatment (rather than to a particular target) by using the breast adenocarcinoma cell line MCF7 as a discriminating cell model. Such a pseudotarget-agnostic strategy identified compounds that inhibited pathways involved in breast cancer survival and also disregarded compounds with low cell penetrability (= deficient drug-likeness). After two rounds of design, synthesis, and screening of highly focused libraries, novel candidates with low micromolar activity in MCF7 cells and high solubility in water were identified (compounds 8d and 9d). Elucidation of the potential target/s linked with their bioactivity was followed by a literature survey to explore chemical motifs that could further improve these phenotypic leads. From that point, two independent medicinal chemistry campaigns were initiated. This manuscript has focused on the development of SRC inhibitor 11a, a novel pyrazolopyrimidine analogue that exhibits potent antiproliferative properties against breast cancer cells, halts MDA-MB-231 cell migration, and inhibits intracellular SRC signaling at low nanomolar concentrations. Remarkably, kinase profiling revealed that 11a is the first small molecule that inhibits SRC at subnanomolar concentration (IC50 < 0.5 nM) with a 1000-fold selectivity over ABL. This is of relevance because the manifestation of cardiac events, especially in elderly patients, is a well-established adverse effect of ABL inhibition.31,32

**Figure 8.** Immunohistochemical analysis of phospho-SRCY416 in human tumor xenografts. (a) Images of representative sections (low and high resolution) of HCT116 xenografts from (left) untreated mice and (right) mice treated with 11a (n = 4). (b) Histoscore analysis (6–7 sections analyzed per experiment). Quantification of immunohistochemistry across tumor tissue sections from untreated animals (water) and 11a treated groups performed in blinded fashion. P value calculated from t-test.
Financial properties, including excellent water solubility and cLogP of 2.2, prompted us to investigate further the biological properties of this compound. Biochemical and in silico studies confirm that 11a is a Type I kinase inhibitor of SRC and provided rationale for its selective affinity for SRC over ABL. Screening of hERG channel and CYP induction, together with in vitro and in vivo PK/PD assays (including zebrafish PD/safety studies; support the nomination of 11a as a SRC-targeting drug candidate that could potentially offer a superior therapeutic window than dual SRC/ABL inhibitors.60,61 This novel SRC-targeting inhibitor belongs to a second generation of inhibitors that do not target the ABL kinase,62 being the first one to demonstrate suppression of SRC phosphorylation in vivo. Finally, in concordance with the primary role of SRC in driving tumor invasion and drug resistance mechanisms,61 more in-depth preclinical proof-of-concept studies using appropriate in vivo models of disease progression and rational drug combination studies will be required to demonstrate the efficacy and therapeutic value of SRC/no-ABL inhibitors in cancer treatment.

**EXPERIMENTAL SECTION**

**Synthetic Procedures. General Methods.** Microwave-irradiated reactions were carried out in a Biotage Initiator microwave synthesizer. Nonmicrowave reactions were performed under an inert atmosphere in a drybox. General methods for microwave reactions were carried out in a Biotage Initiator microwave synthesizer. Nonmicrowave reactions were performed under an inert atmosphere in a drybox. General methods for microwave reactions were carried out in a Biotage Initiator microwave synthesizer.

**Formation of Compound 11a.** To a solution of 1-(2,2-Diethoxyethyl)-3-(1H-pyrrolo[2,3-b]pyridin-5-yl)pyrazolo[3,4-d]pyrimidin-4-amine (8d) in dioxane (10 mL) and water (1 mL) were added 7-azaindole-5-carboxylic acid pinacol ester (1.1 equiv, 642 mg, 4.5 mmol), and the mixture heated in the microwave at 120 °C for 1 h. EtOAc and water (50 mL) were added to the mixture, and the organic layer separated. The aqueous layer was washed with EtOAc (50 mL, x3), and the organics combined and concentrated in vacuo. The crude product was purified by column chromatography.

**Synthesis of Compound 11a.** To a solution of 1-(2,2-Diethoxyethyl)-3-(1H-pyrrolo[2,3-b]pyridin-5-yl)pyrazolo[3,4-d]pyrimidin-4-amine (8d) in dioxane (10 mL) and water (1 mL) were added 7-azaindole-5-carboxylic acid pinacol ester (1.1 equiv, 642 mg, 4.5 mmol), and the mixture heated in the microwave at 120 °C for 1 h. EtOAc and water (50 mL) were added to the mixture, and the organic layer separated. The aqueous layer was washed with EtOAc (50 mL, x3), and the organics combined and concentrated in vacuo. The crude product was purified by column chromatography.

**Synthesis of Compound 6a.** Compound 6a was obtained from 1-(2,2-Diethoxyethyl)-3-(1H-pyrrolo[2,3-b]pyridin-5-yl)pyrazolo[3,4-d]pyrimidin-4-amine (8d) by a similar procedure.

**Formation of Compound 11a.** To a solution of 1-(2,2-Diethoxyethyl)-3-(1H-pyrrolo[2,3-b]pyridin-5-yl)pyrazolo[3,4-d]pyrimidin-4-amine (8d) in dioxane (10 mL) and water (1 mL) were added 7-azaindole-5-carboxylic acid pinacol ester (1.1 equiv, 642 mg, 4.5 mmol), and the mixture heated in the microwave at 120 °C for 1 h. EtOAc and water (50 mL) were added to the mixture, and the organic layer separated. The aqueous layer was washed with EtOAc (50 mL, x3), and the organics combined and concentrated in vacuo. The crude product was purified by column chromatography.

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0.598 mmol, 126.8 mg) was added, and the mixture allowed to stir for 17 h overnight. The mixture was concentrated in vacuo, and the product purified by column chromatography, MeOH/DCM (0–10% then 5–20 drops of NH4aq per 100 mL) to give a light orange/brown solid (163.8 mg, 0.405 mmol, 99%).1H NMR (500 MHz, MeOD) δ 8.22 (s, 1H), 4.49 (t, J = 6.4, 2H), 3.32 (s, 3H), 3.15 (d, J = 12.1, 2H), 2.96 (dd, J = 16.0, 8.0, 4.0, 1H), 2.91 (t, J = 6.4, 2H), 2.72 (s, 6H), 2.15 (td, J = 12.0, 2.0, 2H), 2.04–1.96 (m, 2H), 1.54 (qd, J = 12.2, 3.9, 2H);13C NMR (126 MHz, MeOD) δ 158.07 (C), 155.67 (CH), 153.70 (C), 103.59 (C), 86.97 (C), 63.18 (CH), 55.86 (CH2), 52.91 (CH3), 44.37 (CH2), 39.31 (CH3), 26.42 (CH2); MS (ES +ve) [M + H]+: 416.2. 

**Western Blotting.** Cells were plated at 1 × 105 cells/well in 2 mL of DMEM medium containing 10% FBS and 2 mM L-glutamine in 6-well plates and incubated at 37 °C with 5% CO2. After 24 h, the media was replaced and replaced with 2 mL of DMEM medium containing 0.1% FBS and 2 mM L-glutamine and the cells incubated for a further 24 h. Two μL of compounds dissolved in DMSO at appropriate concentration was then added to each well and plates incubated for 1.5 h. 222 μL of FBS was then added to each well (giving a final concentration of 10%), and cells incubated for 1 h. Cell lysates were then prepared using 100 μL of cell lysis buffer (1% Triton X-100, 50 mM HEPES, pH 7.4, 150 mM NaCl, 1.5 mM MgCl2, 10 mM NaF, 10 mM sodium pyrophosphate, 1 mM Na3VO4, 10% glycerol and protease and phosphatase inhibitors) per well. The total cell protein concentration in each lysate was determined using precision red advanced protein reagent #2 from Cytoskeleton. Lysates were normalized to 0.2 mg/mL and boiled at 100 °C for 3 min in SDS-PAGE sample buffer (40% glycerol, 8% SDS, 0.1 M DTT, 0.25 M Tris-HCl, pH 6.8). Samples were resolved by SDS-PAGE using BioRad 4–15% precast gels over 60 min at 140 V and transferred to PVDF membranes over 150 min at 210 mA. Nonspecific antibody binding was blocked by incubation of membranes for 1 h at room temperature using Western blotting reagent solution (Roche) prior to adding primary antibodies in 0.5% blocking buffer at 4 °C overnight. Membranes were washed with TBS/T (×3, 5 min) and secondary antibody linked to horseradish peroxidase (HRP) added for 1 h at room temperature. Following further washing with TBS/T (×3, 5 min) and TBS (×2, 5 min), HRP was revealed by peroxidase enhanced chemiluminescence (Pied ECL from Roche) and bands visualized using X-ray film or the ChemiDoc MP Imaging System from BioRad.

**Cell Migration Assay.** Cells were plated at 50,000 cells/well in 100 μL of DMEM medium containing 10% FBS and 2 mM L-glutamine in a 96-well ImageLock plate from Essen BioScience and left overnight to adhere in an incubator at 37 °C and 5% CO2. Scratch wounds were created in each well using the WoundMaker supplied by Essen BioScience, and each well washed with media (100 μL) to remove floating cells. 95 mL of fresh media was added in each well. Compounds, including DMSO, were prepared at 20× in DMEM medium in a separate plate, and 5 μL was then added to each well containing cells. Untreated cells were incubated with DMSO (0.1% v/v). Images were recorded every 30 min using the IncuCyte-ZOOM for 24 h. Analysis of cell migration into the wound was performed using the IncuCyte software.

**Kinetin Screening Assays - IC50 Determination.** Radiosotope-based assay ([γ-32P] ATP) consisting of measuring ATP incorporation on the substrate (poly [Glu, Tyr] 4:1) relative to DMSO. Compound IC50 values were determined from 10-point, 1:3 dilution curves starting at 10 μM, by Reaction Biology Corp, with 10 μM ATP.

**Evaluation of ATP Competition and Ki Value for 11a against SRC.** Materials: Inhibitor: 10 mM stock solution of 11a in DMSO. Kinase reaction buffer: 20 mM HEPES-HCl, pH 7.5, 10 mM MgCl2, 1 mM EGTA, 0.02% Brij35, 0.1 mM Na3VO4, 0.02 mg/mL BSA, 2 mM DTT, and 1% DMSO. Kinase: SRC; recombinant human full length kinase domain (accession no. NP_005408), C-terminal His-tagged expressed in insect cells, Msapp = 62.3 kDa. Substrate for kinase: poly [Glu, Tyr] 4:1. Reaction conditions: 0.6 mM SRC, 0.2 mg/mL
poly [Glu, Tyr] 4:1, and varied ATP. Experimental procedure: The kinase assays were performed at room temperature. 11a was added as 10-point dose IC₅₀ mode into the enzyme/substrate mixture using acoustic technology with 20 min preincubation. Then the five concentrations of ATP were added to initiate the reaction. The activity was monitored every 5–15 min for a time course study. The ATP, substrate, and 11a concentrations tested were as follows: (i) ATP concentrations tested: 1, 5, 15, 75, and 150 μM ATP; (ii) substrate concentration: constant at 0.2 mg/mL; (iii) 11a concentrations tested: 10-point dose IC₅₀ with 3-fold serial dilution started at 100 nM; (iv) time points measured: 0, 5, 10, 15, 20, 30, 45, 60, 75, 90, 105, 120 min.

Docking Studies. Crystal structures of SRC and ABL were examined for their suitability for docking. PDB 3GEQ was selected to represent SRC as it contained pyrazolopyrimidine ligand PP2, which most closely resembles compounds PP20 and 11a, and its DFG motif is in the active conformation. Similarly, PDB 4TWP was selected as the ABL structure as its DFG motif is also flipped in the ATP-interacting position. The side chain of Asp381 in the ABL DFG motif was manually rotated in PyMol to match the ATP-binding mode of the equivalent Asp in SRC. ABL in this structure also contains a T316I mutation which was reverted to wild type. Models of PP20 and 11a were built using the structure and binding mode of PP2 as a template. Water molecules and other heteroatoms were removed from the optimized structures, and the program PDB 2PQR 1.865 used to assign position-optimized hydrogen atoms, utilizing the additional PropKa60 algorithm with a pH of 7.4 to predict protonation states. The MGLTools 1.5.4 utility prepare receptor4.py was used to assign Gasteiger charges to atoms. Hydrogen atoms were assigned to compound structures using OpenBabel 2.3.2,67 utilizing the -p option to predict the protonation states of functional groups at pH 7.4. The MGLTools utility prepare_ligand4.py67 was used to assign Gasteiger charges and rotatable bonds. Autodock 4.2.369 was used to automatically dock the compounds into the ATP binding pocket of the crystal structures. A grid box that encompassed the maximum dimensions of the cognate ATP ligand plus 12 Å in each direction was used. The starting translation and orientation of the ligand and the torsion angles of all rotatable bonds were set to those of the modeled compounds. The Autogrid grid point spacing was set at 0.2 Å. The Autodock parameter file specified 10 Lamarckian genetic algorithm runs, 15,000,000 energy evaluations and a population size of 300.

Zebrafish PD/Toxicology Assay. Transgenic Tg(brn3c:mGFP) zebrafish embryos were collected from breeding pairs and reared at 28 °C in E3 embryo media. One dpf embryos were treated with 11a at different doses (10–750 μM) at 20 hpf, 36 hpf and 48 hpf, and DMSO (0.1% v/v). Zebrafish embryos were imaged by fluorescent microscopy at 72 hpf. Safety assays: Wild-type zebrafish embryos were collected from AB-TPL breeding pairs and reared at 28 °C in E3 embryo media. One dpf embryos were treated with 11a or dasatinib at 100 μM, and DMSO (0.1% v/v) as negative control, for 4 h before being washed off and replaced with fresh E3 media. For PP20 treatment, the fish were incubated for 2 h post-amputation then replaced with fresh E3 media. The embryos were left to develop in E3 media at 28 °C for 2 d, after which, they were imaged by light microscopy. Zebrafish husbandry was performed under Home Office License in compliance with the Animals (Scientific Procedures) Act 1986 and approved by the University of Edinburgh Ethics Committee. In Vivo PK Analysis. Three female cluster of differentiation 1 (CD1) mice, 25–30 g, were dosed per administration route per time point, per compound. 11a was administered either orally or intravenously (typical dose level of 10 mg compound per kg body weight). Animals were given free access to food throughout the study. At the following time points, the animals were anaesthetised, blood collected in heparinised tubes, and animals sacrificed: (i) oral dosing: 0.08, 0.25, 0.5, 1, 2, 4, and 8 h post-dose; (ii) IV dosing: 0.08, 0.25, 0.5, 1, 2, 4, and 8 h post-dose.

In Vivo PD Study. Tumor xenographs were generated in mice by injection of 2 million HCT116 cells subcutaneously. Tumors were allowed to grow until 3–4 mm in diameter. A daily dose of 50 mg/kg of 11a in pure water was administered by oral gavage. Mice were sacrificed 3 h after the last dose and tumors excised, fixed in 4% formaldehyde in 0.1 M phosphate buffer (pH 7.2), and embedded in paraffin. Sections were cut using a Reichert-Jung 1150/AutoCut microtome to perform phospho-SRC immunochemistry. Antigen retrieval was performed using heat treatment under pressure in a microwave oven for 10 min in 10 mM citrate buffer pH = 6. Sections were blocked for endogenous peroxidase followed by incubation with antiphospho-SRC antibody (Cell Signaling Technology) (1:200 dilution) at 4 °C overnight. Staining was developed using EnVision (Dako) and diaminobenzidine (Dako) before slides were counterstained in hematoxylin, dehydrated, and mounted in dibutyl phthalate xylene. Slices were imaged on a NanoZoomer digital slide scanner, Hamamatsu. Staining was scored by a single experienced observer, blinded to treatment, using a weighted histoscore method.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jmedchem.6b00065. Synthesis and characterization of compounds 2 to 12a–y, in vitro ADME-Tox methods and supporting Figures S1–S12 and Table S1 (PDF) seven-day time-lapse motion picture of MCF7 cell proliferation under treatment (addition at 48 h) with 100 μM of 8d with Nucview488 (apoptosis fluorescent marker) (AVI) seven-day time-lapse motion picture of MCF7 cell proliferation under treatment (addition at 48 h) with 100 μM of 9d with Nucview488 (apoptosis fluorescent marker) (AVI) Additional compound data (XLSX)

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Notes
The authors declare the following competing financial interest(s): The authors declare that a patent application (No. GB1508747.1) is pending on compounds 11-12.

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ANGES USED
ATP, adenosine triphosphate; CD3OD, deuterated methanol; CD1, cluster of differentiation 1; CML, chronic myeloid
leukemia; CYP, cytochrome P450; DFG, aspartate-phenylalanine-glycine; DMF, dimethylformamide; DMEM, Dulbecco’s modified eagle’s media; DCM, dichloromethane; DMPK, drug metabolism and pharmacokinetics; DMSO, dimethyl sulfoxide; dpd, days post-fertilization; IC\textsubscript{50}, half-maximal effective concentration; FAK, focal adhesion kinase; GFP, green fluorescent protein; HLM, human liver microsomes; IC\textsubscript{50}, half-maximal inhibitory concentration; PD, pharmacodynamics; rt, room temperature; SAR, structure–activity relationship; SFK, SRC family of kinases; SYF, murine embryonic fibroblasts.

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