

Structure-Guided Inhibitor Design Expands the Scope of Analog-Sensitive Kinase Technology

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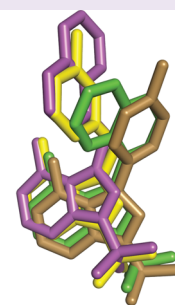
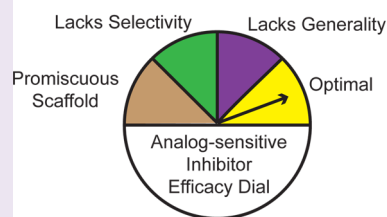
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Supporting Information

ABSTRACT: Engineered analog-sensitive (AS) protein kinases have emerged as powerful tools for dissecting phospho-signaling pathways, for elucidating the cellular function of individual kinases, and for deciphering unanticipated effects of clinical therapeutics. A crucial and necessary feature of this technology is a bioorthogonal small molecule that is innocuous toward native cellular systems but potently inhibits the engineered kinase. In order to generalize this method, we sought a molecule capable of targeting divergent AS-kinases. Here we employ X-ray crystallography and medicinal chemistry to unravel the mechanism of current inhibitors and use these insights to design the most potent, selective, and general AS-kinase inhibitors reported to date. We use large-scale kinase inhibitor profiling to characterize the selectivity of these molecules as well as examine the consequences of potential off-target effects in chemical genetic experiments. The molecules reported here will serve as powerful tools in efforts to extend AS-kinase technology to the entire kinome and the principles discovered may help in the design of other engineered enzyme/ligand pairs.



Eukaryotic protein kinases catalyze the transfer of phosphate from ATP to serine, threonine, or tyrosine residues on substrate proteins. The resulting phosphorylation event constitutes a critical mode of post-translational protein modification and a core mechanism for cellular signal transduction. Nearly every physiological process in eukaryotic cells involves phospho-signaling networks, and many human diseases occur when these pathways are dysregulated.¹ Potent and selective inhibitors of protein kinases are valuable tools for probing these networks and for elucidating the cellular functions of individual kinases.² However, due to the large number of protein kinases in a cell and their highly homologous active sites, it has proven challenging to find specific inhibitors for many individual kinases.³ Our laboratory has developed a chemical genetic technique that enables systematic generation of highly specific inhibitors for individual kinases.⁴ This approach combines the advantages of a small molecule inhibitor (temporal resolution, reversibility, etc.) with the specificity of a genetic manipulation. When a residue at a structurally conserved position in the kinase active site (termed the gatekeeper⁵) is mutated from the natural bulky amino acid side

chain (methionine, leucine, phenylalanine, threonine, etc.) to smaller glycine or alanine, a novel pocket that is not found in wild-type (*WT*) kinases is created within the ATP-binding site. Such an engineered kinase is termed an analog-sensitive (*AS*) allele because it can be potently and specifically targeted by inhibitor analogs that contain a bulky substituent complementing the enlarged ATP binding pocket.⁶

A major goal of our laboratory is to extend AS-kinase technology to the entire kinome; the fact that nearly all kinases contain a bulky gatekeeper residue suggests this undertaking is possible in principle. However, two criteria must be satisfied for a particular kinase-of-interest to be amenable to the approach. First, the kinase must tolerate mutation of the gatekeeper residue to glycine or alanine without severe loss of catalytic activity or cellular function. Second, a potent and bioorthogonal small molecule inhibitor of the AS-kinase must be identified. In the course of our studies we have encountered a number of

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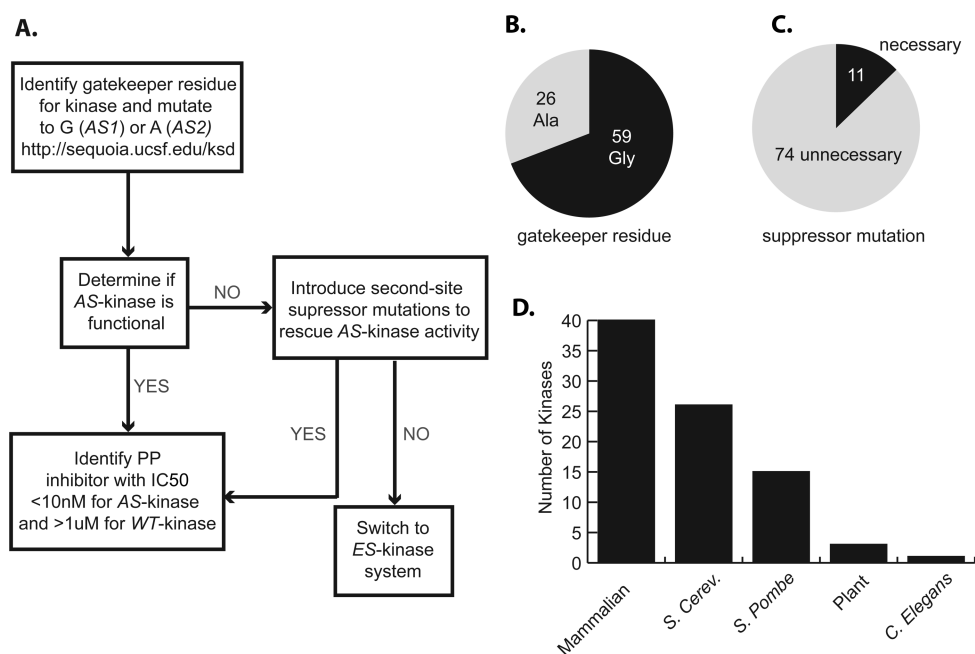


Figure 1. (A) Flow chart outlining a systematic approach for application of AS-kinase technology. (B) The number of reported AS-kinases with alanine, A, or glycine, G, gatekeeper residues. (C) The number of reported AS-kinases that require suppressor mutations to rescue activity. (D) The number of AS-kinases that have been reported in peer-reviewed publications for each class of kinase.

kinases that do not meet one or both of the criteria; they are either intolerant of the gatekeeper mutation or insensitive to available pyrazolo[3,4-*d*]pyrimidine (PP) inhibitors. We have developed a systematic workflow to overcome each of these limitations as they present themselves (Figure 1A). For example, if the kinase does not tolerate a glycine gatekeeper, which provides the maximal expansion of the ATP pocket, the larger amino acid alanine may be used instead (Figure 1B). We have also demonstrated that the introduction of second-site suppressor mutations can be employed to rescue the activity of weakened AS-kinase alleles⁷ (Figure 1C). In extreme cases where second-site suppressor mutations fail to rescue activity, a second approach termed an electrophile-sensitive (*ES*)-kinase may be used.⁸ In *ES*-kinases the gatekeeper is mutated to cysteine instead of alanine or glycine, thereby retaining a more hydrophobic residue at the gatekeeper position, maintaining the integrity of the hydrophobic spine of the kinase,⁹ and thus enhancing enzyme activity, while sensitizing the *ES*-kinase to PP inhibitors bearing an appropriately positioned electrophile. Once an active AS-kinase or *ES*-kinase allele has been engineered, a potent and selective AS-kinase or *ES*-kinase inhibitor must be identified from a small library of PP molecules based on the semi-promiscuous tyrosine kinase inhibitor PP1.

PP1 was initially identified as a potent ATP-competitive inhibitor of Src family kinases.¹⁰ The cocrystal structure of PP1 bound to a Src family kinase, Hck, showed that the pyrazolopyrimidine core of PP1 mimics the adenine ring of ATP in its binding to the nucleobase pocket while the *p*-tolyl group at the C3 position projects into a deep hydrophobic pocket situated between the gatekeeper (T338) and the catalytic lysine residue (K295).¹¹ On the basis of this crystal structure, it was predicted that PP1 analogs with an enlarged C3 substituent would suffer a steric clash with the native gatekeeper residue in *WT*-kinases. However, mutation of the gatekeeper to glycine or alanine should create extra space to

accommodate the enlarged C3 substituent. It has been demonstrated that potent and specific inhibitors can be readily identified for various AS-kinases by screening a small panel of PPs with enlarged C3 substituents.⁴

INA-PP1 and 1NM-PP1 were initially found to be the most effective inhibitors for AS-kinases and have been used to target numerous AS-kinase alleles from a variety of organisms and kinase families (Figure 1D, Supplementary Table S1). In the course of our studies, however, we have encountered a number of AS-kinases that are insensitive to INA-PP1 and 1NM-PP1, thereby precluding the application of our chemical genetic technique. Our initial hypothesis was that the PP structure was suboptimal for some kinases, and thus we sought alternative scaffolds that might be complementary or even superior to PP. For example, we used an inhibitor based on the aminoindazole scaffold (a known Akt binder) to target AS-Akt alleles.^{12,13} However, this molecule and its derivatives were largely inactive toward other AS-kinases (unpublished results). We also attempted to design AS-kinase inhibitors based on purine, pyrimidine, and quinazoline scaffolds,^{14,15} but these molecules also proved to be largely ineffective against diverse AS-kinases. Since the PP scaffold has so far proven to be the most general for AS-kinases, we embarked on a medicinal chemistry effort to find a new PP analog capable of targeting recalcitrant AS-kinases. The result of this undertaking was the identification of the new molecule 3MB-PP1 that has enhanced potency against a diverse set of AS-kinases, including several that were insensitive to all previously known AS-kinase inhibitors.^{12,16–18}

The focus of this study is to understand the mechanism of efficacy of 3MB-PP1 by analysis of cocrystal structures of Src-AS1 kinase bound to 3MB-PP1, INA-PP1, or 1NM-PP1. We then use insights gained from these structures to design next-generation PP inhibitors with increased potency and selectivity for AS-kinases.

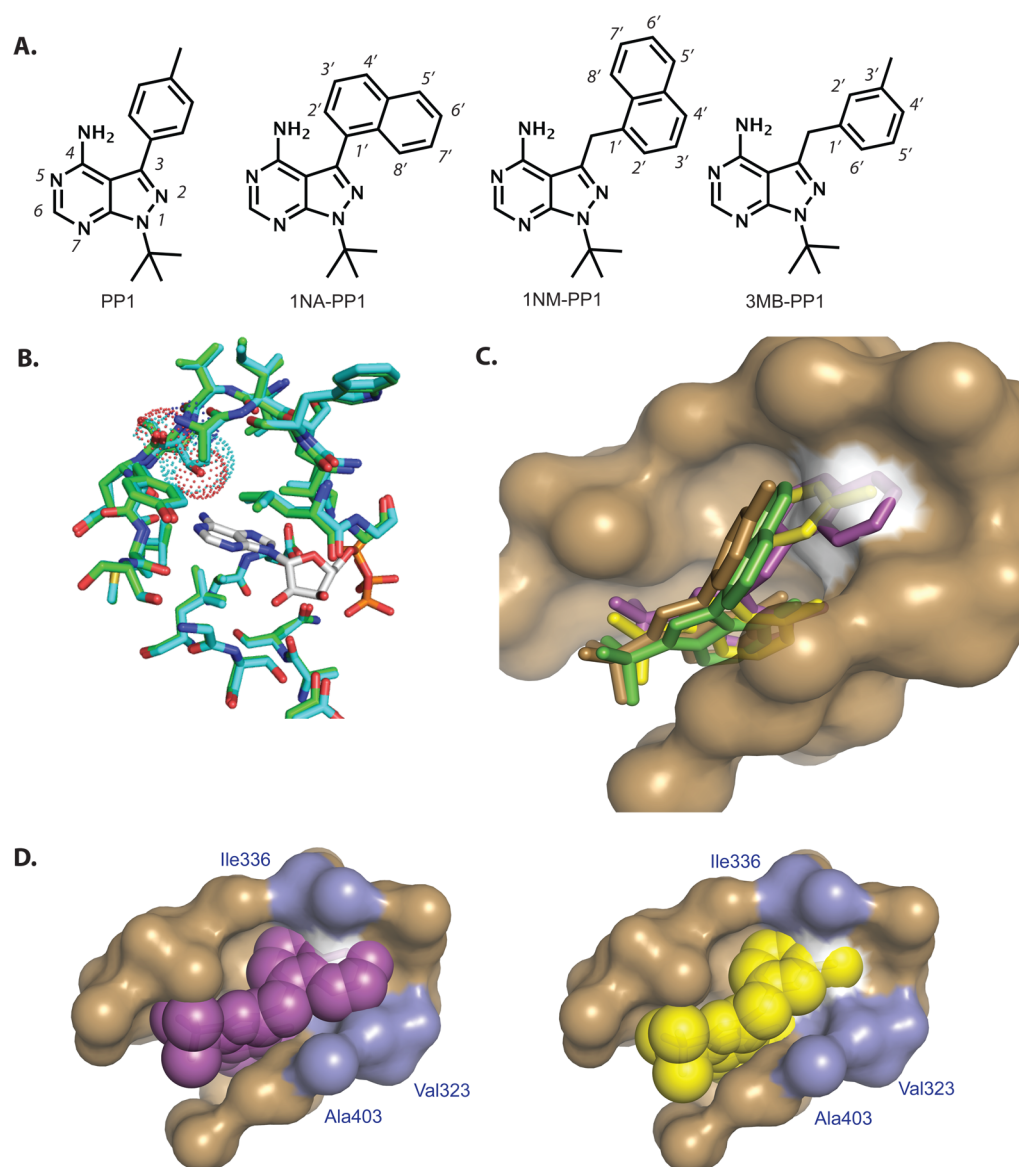


Figure 2. (A) PP1 is a promiscuous kinase inhibitor that targets kinases with medium size gatekeeper residues (Thr, Val, Ser). 1NA-PP1, 1NM-PP1, and 3MB-PP1 are three of the most commonly used AS-kinase inhibitors. (B) Overlay of Src-WT (teal) and Src-AS1 (green) with the WT-gatekeeper surface illustrated in dots and the Src-WT ligand AMP-PNP (white). (C) Overlay of 1NA-PP1 (green), 1NM-PP1 (pink), and 3MB-PP1 (yellow) with the PP1/Hck-WT structure (brown). The surface of the gatekeeper residue is highlighted in white. (D) Overlay of Hck ATP pocket surface with 3MB-PP1 (right, yellow spheres) and 1NM-PP1 (left, pink spheres). Surfaces of residues V323, I336, and A403 are highlighted in blue.

RESULTS AND DISCUSSION

X-ray Crystal Structures of PP Inhibitors Bound to Src-AS1 Reveal Mechanism of Enhanced Efficacy of 3MB-PP1. We previously reported that 3MB-PP1 and a closely related analog **16** (Figure 2A, Figure 3) are capable of targeting several AS-kinases that are impervious to 1NA-PP1 and 1NM-PP1.^{12,16–18} In order to better understand the enhanced efficacy of 3MB-PP1 and the potential deficiencies of 1NA-PP1 and 1NM-PP1 in targeting divergent AS-kinases, we solved cocrystal structures of Src-AS1 (T338G) in complex with 1NA-PP1, 1NM-PP1 or 3MB-PP1 (Figure 2C). Comparison of the Src-AS1 structures to the previously solved Src-WT/AMP-PNP complex reveals little movement of protein backbone atoms (with an RMSD of 0.22 Å) upon introduction of the gatekeeper mutation T338G, indicating that the mutation does not perturb the overall structure of the Src protein (Figure 2B). The most notable effect of the T338G mutation in Src is the expanded

ATP-binding pocket that we hypothesized would accommodate bulky groups at the C3 position of PP1. Next we created an overlay of the Src-AS1 structures with that of the highly homologous tyrosine kinase Hck bound to PP1 (Figure 2C). We found that the PP analogs share nearly identical binding orientations and hydrogen bond interactions with the hinge region of the kinase. As predicted, the bulky naphthyl, naphthylmethyl and 3'-methylbenzyl substituents of 1NA-PP1, 1NM-PP1 and 3MB-PP1, respectively, project toward the gatekeeper and occupy the pocket created by the T338G mutation. While PP1 is accommodated in the ATP pocket of Hck, it is clear that all three AS-kinase inhibitors are prevented from binding due to a steric clash with the Thr gatekeeper residue; however, the degree of clash varies among the compounds. Specifically, the naphthyl ring of 1NA-PP1 (Figure 2C, green) appears to have minimal overlap with the surface of the gatekeeper side chain, while the clash between the

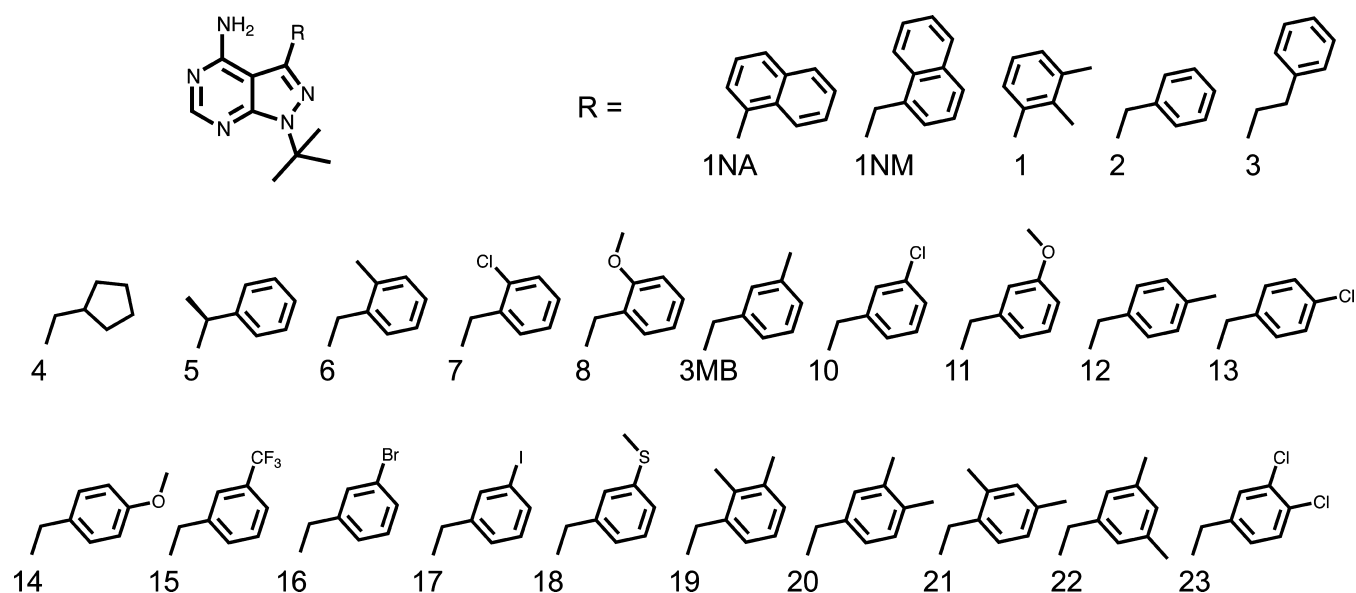


Figure 3. Chemical structures of PP compounds.

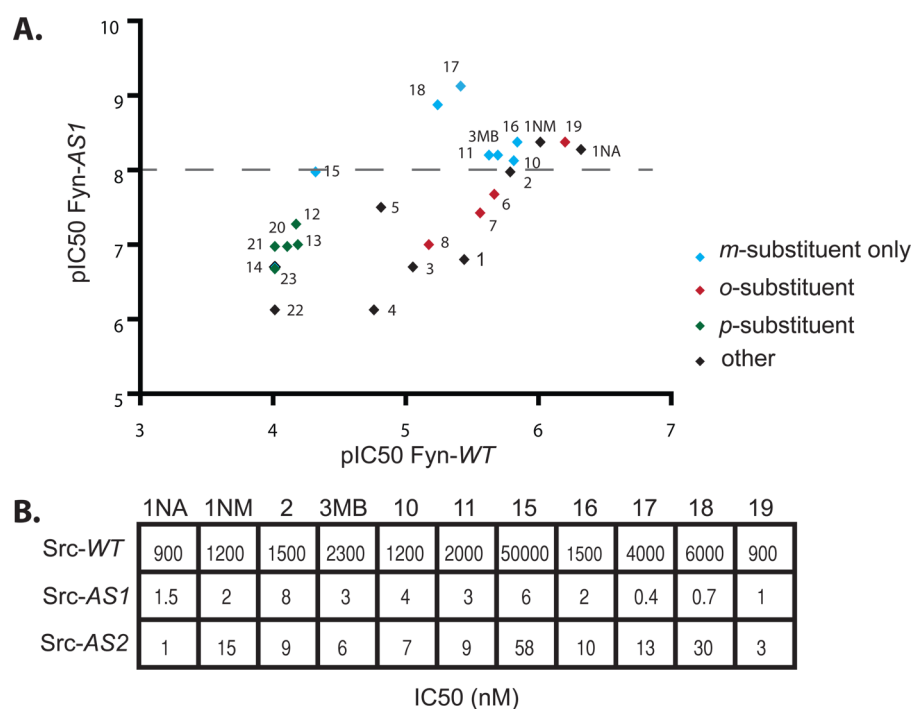


Figure 4. (A) pIC₅₀ of each compound for Fyn-AS1 plotted against the pIC₅₀ for Fyn-WT to illustrate the potency and selectivity of PP inhibitors. The dotted gray line indicates the threshold for compounds with suitable potency (pIC₅₀ > 8) against Fyn-AS1. (B) IC₅₀ values for select PPs against Src-WT, Src-AS1, and Src-AS2. IC₅₀ values were determined from a single dose–response measurement.

gatekeeper and 1NM-PP1 (Figure 2C, purple) is maximal and that with 3MB-PP1 (Figure 2C, yellow) is intermediate. The minimal clash between 1NA-PP1 and a Thr gatekeeper explains why 1NA-PP1 has weak activity toward WT-kinases with small gatekeeper residues but not toward kinases with larger Ile or Phe gatekeeper residues.⁴ In contrast, the C5' position of 1NM-PP1 projects deep into the gatekeeper side chain surface of Hck, thus explaining the enhanced orthogonality of 1NM-PP1 toward WT-kinases relative to 1NA-PP1.

A key feature of both 1NM-PP1 and 3MB-PP1 is the methylene linker between the aryl ring and the heterocyclic core of the PP scaffold. This linkage imbues the molecules with

rotational freedom that allows for optimal orientation of the bulky phenyl and naphthyl rings in the gatekeeper pocket, thereby maximizing potential affinity interactions with the enlarged ATP pocket of AS-kinases. Simultaneously, the methylene linker ensures that the bulky aryl groups project more deeply into the gatekeeper pocket, resulting in greater steric clash with the gatekeeper side chain of WT-kinases. The 3'-methyl group of 3MB-PP1 and C5' of 1NM-PP1 appear to occupy the same position within the ATP binding pocket and provide the most direct steric clash with the gatekeeper side chain. An important difference between 3MB-PP1 and 1NM-PP1, however, is that the additional atoms of the naphthyl ring

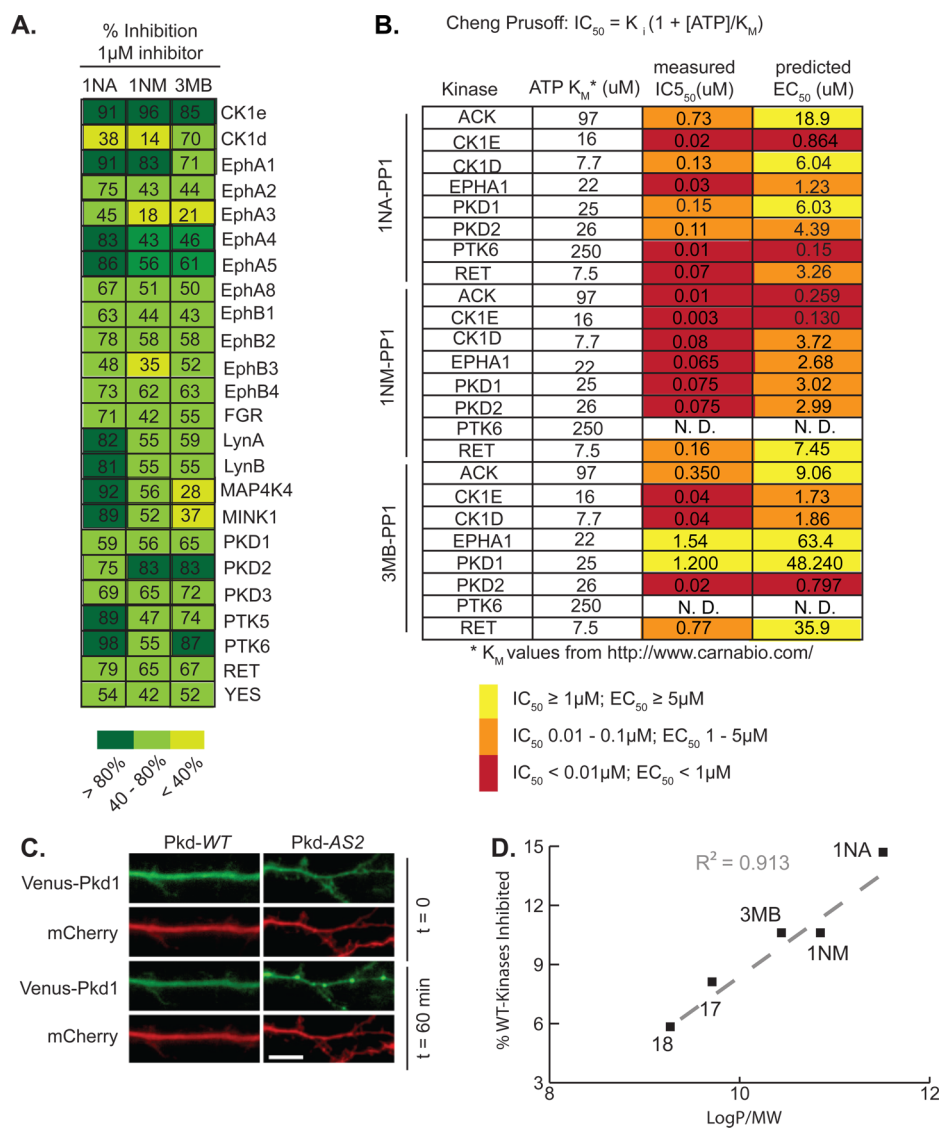


Figure 5. (A) Fraction of activity of wild-type PP-sensitive kinases inhibited by 1 μ M 1NA-PP1, 1NM-PP1, and 3MB-PP1. (B) IC_{50} values were measured for 1NA-PP1, 1NM-PP1, and 3MB-PP1 toward eight WT-kinases. The Cheng–Prusoff equation and K_M values of each kinase for ATP were used to calculate predicted EC_{50} values for 1NA-PP1, 1NM-PP1, and 3MB-PP1 toward eight WT-kinases. (C) Neurons were transfected with Venus-Pkd1-AS2, treated with 1 μ M 1NA-PP1 and analyzed for puncta formation. (D) The fraction of WT-kinases inhibited by 40% or more was plotted as a function of CLogP value for 1NA-PP1, 1NM-PP1, 3MB-PP1, 17, and 18.

of 1NM-PP1 (C6', C7', and C8') provide additional clash with the Hck gatekeeper but also appear to overlap with the surface of other residues in the ATP binding pocket (V323, I336, and A403; Figure 2D). We hypothesize that subtle differences in the ATP binding sites of divergent kinases might alleviate or escalate this undesired interaction and, in extreme cases, prevent 1NM-PP1 from binding to the AS-kinase despite the enlarged gatekeeper pocket. In fact, it has been demonstrated that some kinases bearing larger amino acids at the position analogous to Ala403 can be further sensitized to PP derivatives by mutation of that position to alanine.¹⁹

Structure-Guided Inhibitor Design Yields More Potent, Selective, and General AS-Kinase Inhibitors. Next we sought to test our hypothesis that the benzyl moiety is the optimal orthogonality element for PP AS-kinase inhibitors as well as to design more potent, selective, and general AS-kinase inhibitors using our enhanced understanding of 3MB-PP1 as a guide. Thus, we synthesized a panel of PP derivatives

based largely on the benzyl PP scaffold with various substituents designed to project into the gatekeeper pocket, as does the methyl group of 3MB-PP1 (Figure 3). The PP analogs were synthesized in four steps using a previously described route⁶ and were analyzed for potency and selectivity toward Fyn-AS1 over Fyn-WT. We chose Fyn, a Src family tyrosine kinase, because it is a potent target of PP1;⁵ thus, an inhibitor of AS-kinases that does not target WT-Fyn would ensure that potency was not gained at the expense of selectivity. We plotted the negative logarithm of the IC_{50} value (pIC_{50}) of each compound for Fyn-AS1 against the pIC_{50} for Fyn-WT to illustrate the potency and selectivity of the inhibitors (Figure 4A). Strikingly, all of the new PPs with pIC_{50} (Fyn-AS1) ≥ 8 contain the *meta*-substituted benzyl functional group. Increasing the linkage between the aryl group and the heterocyclic core from one methylene to two (3) or removing the methylene group altogether (1) results in a dramatic decrease in activity. All compounds that are monosubstituted at the C3'

position of the benzyl group are quite potent and selective inhibitors of Fyn-*AS1*. C2'-substituted benzyl PP inhibitors (6–8) showed decreasing potency with increasing size of the substituent, indicating an unfavorable steric clash that can be minimized with smaller substituents. Molecules with substituents at the C4' position (12–14) display an even more dramatic decrease in potency toward both Fyn alleles. Taken together, these results confirm that the C3'-substituted benzyl PP structure is an ideal scaffold for AS-kinase inhibitors.

Notably, two C3'-benzyl PPs, 17 and 18, stand out as being the most potent and selective inhibitors of Fyn-*AS1*; these molecules contain larger, more polarizable groups relative to 3MB-PP1. We wished to verify that the enhanced efficacy of these compounds was a general phenomenon and not unique to Fyn, so we measured their ability to target Src-*WT*, Src-*AS1* (SrcT338G), and Src-*AS2* (SrcT338A). Once again, 17 and 18 have poor ($IC_{50} > 10 \mu\text{M}$) activity toward the *WT*-kinase but are the most potent inhibitors for the kinase bearing a glycine gatekeeper. Compounds 17 and 18 are less potent toward Src-*AS2*, suggesting the iodo and thiomethyl substituents may ideally complement the space created by a glycine gatekeeper but are perhaps too large to fit in the smaller ATP pocket created by the alanine gatekeeper. Taken together, our results suggest that the C3'-substituted PP core is the optimal scaffold for AS-kinase inhibitors; however, the substituent that best complements a particular AS-kinase might vary.

Selectivity Profiles of PP Inhibitors. A crucial property that an effective AS-kinase inhibitor must possess is selectivity; it must potentially inhibit the activity of an AS-kinase but not affect *WT*-kinases or other endogenous proteins. We and others have previously reported that 1NA-PP1 and 1NM-PP1 are very selective but imperfect AS-kinase inhibitors, as they weakly target some *WT*-kinases.^{4,20} To gain a more comprehensive understanding of the off-target effects of our AS-kinase inhibitors we tested the ability of 1 μM 1NA-PP1, 1NM-PP1, 3MB-PP1, 17, and 18 to inhibit the activity of several hundred *WT*-kinases *in vitro* and found that several are indeed sensitive to one or more PP inhibitors (Supplementary Table S2). This is a very high standard for selectivity as AS-kinase inhibitors typically have IC_{50} values $<100 \text{ nM}$ toward AS-kinases and the cellular concentration of the competitor ligand ATP is relatively high (1–10 mM). Nonetheless, we wished to determine if off-target inhibition of *WT*-kinases by PP compounds is significant in the broad panel of AS-technology. We asked which of the *WT*-kinases targeted by our PP inhibitors are likely to be inhibited in cells at concentrations typically used in chemical genetic experiments. To this end we first measured IC_{50} values for 1NA-PP1, 1NM-PP1, and 3MB-PP1 against a representative subset of the PP-sensitive *WT*-kinases identified in our screen (Figure 5A and B). We found that kinases inhibited greater than 80% in our screen generally have IC_{50} values less than 100 nM, while those inhibited between 40% and 80% have IC_{50} values ranging between 100 nM and 1.5 μM (Figure 5B). The correlation between the fractional kinase activity determined in our screen and the measured IC_{50} values gives us confidence that data from the screen can be used to accurately predict the *in vitro* potency of our inhibitors against a large portion of the kinome.

To address whether PP-sensitive *WT*-kinases are likely to be bona fide targets in cells, we calculated the predicted EC_{50} values for each inhibitor/kinase pair using the Cheng–Prusoff equation (Figure 5B), which relates the measured IC_{50} value, the K_M of the enzyme for ATP, and the cellular concentration

of ATP, which we assume to be $\sim 5 \text{ mM}$.^{21,22} PP inhibitors are typically used in mammalian cells at concentrations between 500 nM and 5 μM (Supplementary Table S1). We calculated that many of the identified PP-sensitive *WT*-kinases would only be targeted in cells at concentrations greater than 5 μM , and we therefore predicted their cellular inhibition should be minimal under typical experimental conditions.

We sought to confirm that weak *in vitro* inhibition of *WT*-kinases by PP molecules is not a practical impediment to our technique by demonstrating the analog-sensitive approach can be applied to Protein Kinase D1 (Pkd1), one of the few kinases inhibited by all PP analogs tested. This unusual sensitivity to PP analogs may be due to a lysine residue at position 667 that occurs in all three Pkd family members in place of a conserved hydrophobic amino acid found in nearly all other kinases. K667 is located in the hinge region of the kinase domain, and it is possible that this amino acid side chain allows greater flexibility in the inhibitor binding orientation, although this remains to be tested. Pkd1 is a widely expressed Ser/Thr protein kinase involved in a number of physiological processes including the trafficking of membrane proteins in neurons.²³ We transfected HEK 293T cells with Pkd1-*AS2* (M665A) or Pkd1-*WT* and found that a concentration of 1 μM 1NA-PP1 completely abrogates autophosphorylation by the analog-sensitive allele but not the wild type kinase, despite the fact that Pkd1-*WT* is inhibited by 1NA-PP1 *in vitro* with an IC_{50} of 150 nM (Supplementary Figure S1). We wished to examine the effects of 1NA-PP1 on a physiological function of Pkd1, and thus we examined the localization of the Pkd1-*WT* and Pkd1-*AS2* in the presence of inhibitor. Kinase-dead alleles of Pkd1 have been shown to constitutively localize to the *trans*-Golgi network (TGN) and TGN-derived vesicles.²⁴ We coexpressed Venus-tagged Pkd1-*WT* and Pkd1-*AS2* in neurons, assessed puncta formation in the presence of 1 μM 1NA-PP1, and found that Venus-Pkd1-*WT* neurons were unaffected, while Venus-Pkd1-*AS2* recapitulated the kinase-dead phenotype, forming discrete puncta after 1 h of inhibitor treatment (Figure 5C).

The example of Pkd1 demonstrates that kinases predicted to be targeted by PP inhibitors in cells with EC_{50} values greater than several micromolar are unlikely to be inhibited under the typical conditions of chemical genetic experiments. Additionally, 1NA-PP1 and 3MB-PP1 have been used to selectively target AS-alleles of Ret, EphB2, EphB3, and EphB4 in cells and mice despite weak inhibition of their *WT*-alleles *in vitro*.^{25,26} Thus, many of the *WT*-kinases identified as targets of 1NA-PP1, 1NM-PP1, or 3MB-PP1 in our screen and in the work of others are not likely to be bona fide targets in a cellular context.

As discussed above, weak *in vitro* inhibition of *WT*-kinases by PP inhibitors is unlikely to be relevant in cells due to competition with the high concentration of ATP. However, we postulate that such weak inhibition might serve as a proxy measurement for overall promiscuity of molecules toward unrelated proteins in a cell. Specifically, PP inhibitors that bind to fewer *WT*-kinases are less likely to bind to diverse proteins including metabolic enzymes capable of inactivating drug molecules *in vivo*. An important physical property that contributes to the promiscuity of small molecule inhibitors is lipophilicity; lipophilic molecules tend to bind nonspecifically to hydrophobic regions of proteins such as ligand binding sites.²⁷ Furthermore, lipophilic molecules are more likely to have poor bioavailability, poor solubility and to form aggregates that may nonspecifically inhibit unrelated enzymes.²⁸ We found that the promiscuity of our PP inhibitors was related to both

the lipophilicity and the molecular weight of the molecule; larger molecules (greater molecular weight) are less promiscuous, while more lipophilic molecules (greater LogP) tend to be more promiscuous. We plotted the fraction of *WT*-kinases whose activity is inhibited by 40% or more as a function of LogP/MW for 1NA-PP1, 1NM-PP1, 3MB-PP1, **17**, and **18** and discovered this yields a linear relationship (Figure 5D). This observation was surprising as we initially expected 1NM-PP1 to be the most selective inhibitor due to the greater steric clash between the naphthyl moiety and the ATP-binding pocket observed in the overlay of the 1NM-PP1/*Src-AS1* and the Hck structures. In fact 3MB-PP1 has a selectivity profile that is similar to that of 1NM-PP1, while **17** and **18** inhibit a much smaller fraction of *WT*-kinases. We interpret these results to reflect the fact that two mechanisms of selectivity are at work. Increasing size of the functional group that projects toward the gatekeeper contributes to selectivity as we originally designed. However, minimizing lipophilicity also contributes by reducing nonspecific affinity of the molecules for the hydrophobic ATP binding pocket of kinases. This discovery highlights the importance of a comprehensive evaluation of drug molecules that incorporates consideration of physical properties such as lipophilicity in addition to the more obvious metrics of on-target potency (IC_{50} , EC_{50} , etc).

Summary. We have reported the mechanism of enhanced efficacy of the AS-kinase inhibitor 3MB-PP1 as well as the design and development of new, more potent, and selective AS-kinase inhibitors. We employed X-ray crystallography to decipher the binding modes of 1NA-PP1, 1NM-PP1, and 3MB-PP1 and discovered that the methylene linker of benzyl PP analogs projects the phenyl group toward the gatekeeper, preventing binding to most *WT*-kinases while providing rotational flexibility that allows optimal orientation to complement the enlarged ATP pocket of AS-kinases. We used insights gained from these structures to design a small panel of benzyl-PPs and discovered two molecules, **17** and **18**, that are the most potent and selective AS-kinase inhibitors yet developed. The increased potency and reduced off-target effects of the new PPs combine to yield a major advantage when targeting an individual AS-kinase in cells or organisms that express hundreds of native kinases. Recent studies have identified 3MB-PP1 and **16** as capable of targeting at least 10 AS-kinases that are resistant to inhibition by any other known PP analog.^{12,16–18} Importantly, these kinases are from several different organisms and span four of the seven major kinase families (CMGC, STE, AGC, and TK), suggesting the enhanced efficacy of the inhibitors is not unique to the tyrosine kinases described in this report. We also examined the potential limitations presented by off-target inhibition of *WT*-kinases by PP inhibitors and discovered that many of the *WT*-kinases inhibited *in vitro* are unlikely to be targeted under cellular conditions. We have combined 1NA-PP1 and 1NM-PP1 with a panel of 3-substituted benzyl PP molecules to create a repertoire of potent and selective AS-kinase inhibitors that allows us to extend AS-kinase technology to the edges of the kinome and propels us toward the larger goal of understanding the logic of cellular signal transduction.

METHODS

Chemical Synthesis. Materials obtained commercially were reagent grade and were used without further purification. ¹H NMR and ¹³C NMR spectra were recorded on a Varian 400 spectrometer at 400 and 100 MHz, respectively. High-resolution electron impact mass

spectra were recorded on a MicoMass VG70E spectrometer at the University of California-San Francisco Center for Mass Spectrometry. Reactions were monitored by thin layer chromatography (TLC), using Merck silica gel 60 F₂₅₄ glass plates (0.25 mm thick). Flash chromatography was conducted with Merck silica gel 60 (230–400 mesh).

Protein Crystallization. *Src-AS1* Cocrystal Structure with 1NA-PP1. The structure of 1NA-PP1 in complex with *Src-AS1* (residues 83–533, including the SH3, SH2, kinase, and tail regions) was determined essentially as previously described for an N6-benzyl-ADP complex.²⁹ Briefly, the *Src-AS1* protein, homogeneously phosphorylated on Tyr 527, was concentrated to 10–15 mg mL⁻¹ in storage buffer (20 mM HEPES [pH 7.6], 100 mM NaCl, and 10 mM DTT) and combined with a slight molar excess of 1NA-PP1. The complex was crystallized in hanging drops at 22 °C by combining 2 μL of the above solution with 1 μL of a well solution (50 mM PIPES [pH 6.5], 12% PEG4000, 10 mM DTT). Crystals were briefly transferred to a buffer containing crystallization solution plus 20% glycerol and frozen at –165 °C. Diffraction data were recorded with a Princeton 2k CCD detector on the A1 beamline at CHESS. All diffraction data were integrated and scaled with the programs DENZO and SCALEPACK. The structure was determined by molecular replacement using PDB entry 2SRC as a search model and refined using Refmac. Data collection and refinement statistics are available in Supplementary Table S3.

Src-AS1 Cocrystal Structures with 1NM-PP1 and 3MB-PP1. The kinase domain of *c-Src-AS1* was expressed and purified as previously described.¹⁵ *Src*-drug complexes were formed by mixing 10 mM DMSO stocks of the drug compounds into 100 μM *Src*, 100 mM NaCl, 50 mM Tris [8.0], 2 mM DTT, and 5% glycerol at RT. Crystals were obtained in hanging drops of a 1:1 mixture (protein-drug and mother liquor) by vapor diffusion atop a mother liquor solution of 100 mM MES [6.5], 50 mM NaAc, 12% glycerol, and 6% PEG 4000. Just prior to data collection, crystals were flash frozen in mother liquor containing 25% glycerol. Data for the 1NM-PP1 and 3MB-PP1 structures were collected at the Advanced Light Source, beamlines 8.2.2 and 8.2.1, respectively. Data were processed using HKL2000. Molecular replacement solutions for the 1NM-PP1 and 3MB-PP1 structures were solved using Phaser in the CCP4 program suite with chain A of *c-Src* (PDB ID: 1YOJ) as the search model. Structures were optimized through iterative rounds of manual fitting in Coot and using maximum likelihood refinement protocols in Refmac. Coordinate and topology files for the ligands 1NM and 3MB were derived in PRODRG (Schuettelkopf and van Aalten, 2004). Data collection and refinement statistics are available in Table S3.

ASSOCIATED CONTENT

Supporting Information

Compiled list of each reported analog-sensitive kinase allele and inhibitor, inhibitor profiling data, X-ray crystallography data collection and refinement statistics, Western blot demonstrating effect of 1NA-PP1 on Pkd1 autophosphorylation, chemical characterization and kinase assay conditions. This material is available free of charge via the Internet at <http://pubs.acs.org>.

Accession Codes

The atomic coordinates and structure factors of *Src-AS1* in complex with inhibitors have been deposited in the Protein Data Bank with accession codes 4K11 (*Src-AS1*/1NA-PP1), 4LGH (*Src-AS1*/1NM-PP1), and 4LGG (*Src-AS1*/3MB-PP1).

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Notes

The authors declare no competing financial interest.

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