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2 Tipping the Balance of the Protein Kinase PDK1 Allosteric Regulatory Mechanism by Low-  
3 Molecular-Weight Ligands  
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## Summary

Here we describe compounds that bind to the active site of a protein kinase and influence the dynamics and function of a distant regulatory site. AGC protein kinases have a conserved allosteric site, the PIF-pocket, which regulates protein activity and interaction with substrates. The binding of substrate-derived docking peptides and small molecules to the PIF-pocket activates AGC kinases by allosterically affecting the ATP-binding site. In this study, we identify small compounds that bind to the ATP-binding site and affect the PIF-pocket of the kinase PDK1 and describe the molecular details of the mechanism whereby the compounds enhance or inhibit the function of the regulatory site. PDK1 and Aurora kinase inhibitors that bind to the ATP-binding site unintentionally differentially modulate physiological interactions at the PIF-pocket site. The rationale presented here has implications for the rational development of new allosteric drugs.

## Introduction

Allostery is a fundamental and widespread mechanism of intra-molecular signal transmission whereby local perturbations on a protein affect the structure and dynamics of specific distal regions (Changeux, 2012; Goodey and Benkovic, 2008; Nussinov and Tsai, 2013). The term “allostery” that was once restricted to oligomeric proteins is now also used to describe the conformational changes that intramolecularly link two given distant sites on a protein (e.g. an orthosteric site and an allosteric site). The transmission of signals across long distances relies on dynamic coupling between different structural motifs and can be described as a shift in the population or dynamics of conformers (Bray and Duke, 2004). The formulations for the allosteric conformational changes have been studied over the last 50 years (Fenton, 2008; Monod et al., 1965; Nussinov and Tsai, 2014; Weber, 1972). Allostery thus mediates the responses of regulatory proteins to different stimuli along the transduction of cellular signals.

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Although the allosteric process is intrinsically bidirectional, cell signaling most often uses  
allostery unidirectionally, to transduce a signal downstream a signaling pathway (Kuriyan and  
Eisenberg, 2007). Protein kinases as well as other regulatory proteins that participate in  
signal transduction have evolved stringent switches (allosteric sites) that control their  
activation, i.e. modulate the conformation of the ATP-binding site (orthosteric site) in  
response to the appropriate upstream signals. In recent years there is growing interest in the  
development of allosteric drugs (Conn et al., 2014; Fang et al., 2013; Gray and Fabbro,  
2014). However, the development of allosteric drugs is hampered by the poor knowledge of  
the molecular details of the allosteric process and how this can be rationally modulated at will  
with small compounds. Here, we describe the “reverse” allosteric regulation by small  
compounds on the phosphoinositide-dependent protein kinase 1 (PDK1), i.e., how binding of  
molecules to the ATP-binding site (orthosteric site) affects a regulatory allosteric site,  
reversing the direction in which the allosteric regulation between the two sites is observed in  
cell signaling.

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The protein kinase domain is formed by two lobes with the ATP-binding site located in  
the cleft between the two (Fig. 1A) (Zheng et al., 1993). It is currently proposed that  
dynamics is the underlying mechanism for allosteric regulation in protein kinases (Kornev  
and Taylor, 2015). In the prototype protein kinase PKA, the catalytic domain is constitutively  
active, and regulation of the enzymatic activity is provided by interaction with regulatory  
subunits (Taylor et al., 2012). In contrast, other members of the AGC group of protein  
kinases are not constitutively active and use a conserved hydrophobic pocket called the PIF-  
pocket as a key allosteric regulatory site. The PIF-pocket was originally described in the  
PDK1 as a regulatory site that binds PIF (PDK1 Interacting Fragment), a polypeptide  
sequence derived from a PDK1 substrate (Biondi et al., 2000). The pocket is formed by the  
conserved helix  $\alpha$ C,  $\beta$ -strands 4 and 5 and helix  $\alpha$ B (Fig. 1B). The PIF sequence comprises  
an extended hydrophobic motif (HM) present in AGC kinases but has a phosphomimetic Asp  
residue instead of the Ser/Thr phosphorylation site. Phospho-HM polypeptides and PIFtide

1 activate PDK1 and diverse AGC kinases (Biondi et al., 2000; Engel et al., 2006; Frodin et al.,  
2 2002; Yang et al., 2002) *in vitro* by interaction with the PIF-pocket (Fig. S1A,B). In PDK1, the  
3 PIF-pocket plays an additional role in the specific recognition of a subgroup of its substrates.  
4 The HM of PDK1 substrates such as SGK, S6K, PKC, RSK, but not PKB/Akt, require the  
5 docking of their HM to the PIF-pocket of PDK1 to become efficiently phosphorylated  
6 (Bayascas, 2008; Biondi et al., 2001; Collins et al., 2003) (Fig. S1C). We previously  
7 characterized the mechanism of activation of PDK1 upon binding of peptides or small  
8 molecules to the PIF-pocket (Engel et al., 2006; Hindie et al., 2009; Sadowsky et al., 2011;  
9 Stockman et al., 2009; Wei et al., 2010) (Fig. S1D,E). Crystal structures of PDK1 with the  
10 bound reversible activators PS48 (Hindie et al., 2009) and PS210 (Busschots et al., 2012)  
11 revealed structural aspects of the allosteric activation mechanism and the associated  
12 conformational changes. For instance, the crystal structure of PDK1 in complex with PS210  
13 shows the complete closure of the ATP-binding site (Busschots et al., 2012).

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31 The allosteric regulation of protein kinases mediated by the small lobe of the kinase  
32 domain is not restricted to AGC kinases. Aurora kinases are activated by the polypeptides of  
33 their interacting partners, which bind to a site equivalent to the PIF-pocket (Bayliss et al.,  
34 2003). Furthermore, EGF-receptor kinases are activated by dimerization; an activating  
35 kinase interacts with a receiver kinase at a site equivalent to the PIF-pocket (Zhang et al.,  
36 2006). Abl kinase is both allosterically inhibited and activated by intramolecular interaction  
37 with its SH2 domain (Wojcik et al.), which, in the active state, docks at the top of the small  
38 lobe (Dolker et al., 2014; Nagar et al., 2003). However, except for the case of AGC kinases,  
39 these regulatory features mediated by the small lobe have not been investigated using small  
40 compounds.  
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53 In spite of the large interest in drug development to protein kinases over the last 20  
54 years, the allosteric effects of small compounds binding to the ATP-binding site have  
55 remained mostly unexplored. In the present work we investigate the bi-directional allosteric  
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1 regulation of PDK1 using small compounds that bind to the PIF-pocket and ATP-binding site.  
2 The results show how different small compounds that bind at the ATP-binding site produce  
3 allosteric effects on the PIF-pocket regulatory site, enhancing or inhibiting the binding to  
4 PIFtide. This work highlights how the “old” concept of allostery provides new exciting  
5 opportunities for drug development to protein kinases.  
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## 10 11 12 13 14 15 16 17 **Results**

### 18 19 20 **Identification of small compounds that bind to the ATP-binding site and allosterically** 21 **affect the binding of PIFtide to the PIF-pocket of PDK1**

22 PDK1 interacts with the HM-polypeptide PIFtide with high affinity, and this interaction  
23 increases the specific activity of PDK1 *in vitro* as measured by its increased ability to  
24 phosphorylate T308tide, a polypeptide derived from the activation loop of PKB/Akt (Biondi et  
25 al., 2000) (Fig. S1A). The interaction between His-PDK1 and biotin-PIFtide can be measured  
26 by means of alphascreen technology (Fig. 1C) (Busschots et al., 2012). Using this  
27 homogeneous assay, we screened a library of 14400 small molecules (average mol. weight  
28 320 Da) for their ability to affect the PDK1-PIFtide interaction. We identified small  
29 compounds that displaced the interaction, e.g., PSE10 and PS653 (Fig. 1D,E). PSE10 (2-  
30 oxopropyl N-(4-chlorophenyl)-[(2-chloro-6-fluorobenzoyl)amino]methanimidothioate) has two  
31 ring systems joined by a linker and a side chain, resembling compound 1 (Engel et al., 2006),  
32 PS48 (Hindie et al., 2009) and PS210 (Busschots et al., 2012), which we previously  
33 characterized as activators that bind to the PIF-pocket of PDK1. In contrast, PS653 (1,6-  
34 dihydrodibenzo[cd, g]indazol-6-one) presents a small, planar, anthrone-derived structure. We  
35 tested the ability of the newly identified compounds to affect the intrinsic kinase activity of  
36 PDK1 (Fig. 1F). PSE10 activated PDK1 but did not affect the activity of PDK1 with mutations  
37 at the PIF-pocket (Fig. 1F-J). The crystal structure of PSE10 in complex with PDK1<sub>50-359</sub>  
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1 confirmed that PSE10 bound to the PIF-pocket (Fig. S2 and Table S1) in a manner similar to  
2 the binding modes of PS48 and PS210. In contrast, PS653 inhibited the activity of different  
3 PDK1 constructs (Fig. 1G,H; Fig. S3). We previously showed that PDK1 Leu155Glu, which  
4 contains a mutation at the center of the PIF-pocket, is not activated by PIFtide (Biondi et al.,  
5 2000) or by small compounds that bind to the PIF-pocket (Engel et al., 2006). Similarly, other  
6 mutations in the PIF-pocket also partially abolished the activating effect of PIFtide and small  
7 compounds (Engel et al., 2006). In contrast to PSE10, PS653 still inhibited PDK1  
8 [Leu155Glu] and PDK1 [Val127Leu], which have mutations at different key residues in the  
9 PIF-pocket (Fig. 1I,J). These results were not compatible with PS653 binding to the PIF-  
10 pocket; instead, they suggested that the binding site of PS653 was allosterically coupled to  
11 the PIF-pocket.  
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25 The high-resolution crystal structure of PDK1<sub>50-359</sub> in complex with PS653 (Fig. 2A;  
26 Fig. S2C; Table S1) revealed that PS653 binds at the ATP-binding site, thus confirming that  
27 the displacement of PIFtide by PS653 is mediated by a reverse allosteric effect and not due  
28 to a direct competition for the PIF-binding pocket. Like the adenine moiety of ATP and most  
29 ATP-competitive inhibitors, PS653 forms two hydrogen bonds with the main chain atoms  
30 Ser160-CO and Ala162-N of the so-called linker region and is sandwiched in between the  
31 hydrophobic side chains of Leu88, Val96, and Leu212. However, a major difference is that  
32 PS653 extends outside of the usual ATP-binding cavity. Its outer ring is in close contact (3.25  
33 and 3.0 Å, respectively) with the carbonyl groups of both Leu99 of the small lobe and Ala162  
34 of the large lobe.  
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48 We then investigated whether adenine or adenosine, which bind to the same site as  
49 PS653 (Fig. 2B,C; Table S1), could produce similar allosteric effects. Surprisingly, adenosine  
50 *enhanced* the binding of PIFtide to PDK1, providing evidence of cooperativity, whereas  
51 adenine did not produce any allosteric effect on the PIF-pocket (Fig. 2D). Interestingly, the  
52 enhancing effect was observed selectively with adenosine and not with adenine, AMP, ADP,  
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1 ATP, the non-hydrolyzable ATP analog thio-ATP or cAMP (Fig. S3B). The high-resolution  
2 crystal structure of PDK1 in complex with adenosine showed that the ribose moiety forms  
3 hydrogen bonds with Glu166 of the large lobe and an indirect hydrogen bond via a water  
4 molecule with Ser94 of the small lobe (Fig. 2C). Thus, the presence of the sugar moiety in  
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6 between the lobes played a role in the allosteric enhancement of the binding of PIFtide to the  
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8 PIF-pocket. The differential effects caused by distinct compounds binding to the ATP-binding  
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10 site showed that the resulting allosteric effects were highly selective.  
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16 The co-crystal structures revealed the precise binding mode of the allosteric and non-  
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18 allosteric compounds under investigation. The crystal structures, on the other hand, did not  
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20 reveal significant structural differences that could explain the observed allosteric effects.  
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### 23 **Molecular dynamics simulations describe changes to the conformational dynamics of** 24 **PDK1 upon ligand binding**

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26 To obtain more detailed information on the allosteric mechanisms, we analyzed the changes  
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28 in the dynamics of PDK1 upon the binding of the different effectors using atomistic molecular  
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30 dynamic (MD) simulations. We first performed long MD simulations to observe the effect of  
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32 the different ligands on the sub- $\mu$ s dynamics of PDK1. This is an unbiased approach to  
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34 evaluate the whole dynamics of the protein. Interestingly, we identified an equilibrium  
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36 between two populations of interconverting conformations, which differed in the stability of  
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38 the small  $\alpha$ B helix and could be appropriately quantified by the distance between the Ca  
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40 atoms of Lys115 and Lys123 (end-to-end distance), shown as yellow spheres in figure  
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47 S4A,B.  
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49 To fully sample the conformational changes, that in protein kinases can take place on  
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51 timescales ( $\mu$ s to ms) not easily accessible to conventional MD (Saladino and Gervasio,  
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53 2012), and to reconstruct the conformational free energy landscape, we performed long  
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55 multiple-replica Parallel Tempering (Sugita and Okamoto, 1999) simulations in the Well  
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57 Tempered Ensemble (PT-WTE) (Bonomi and Parrinello, 2010). Using this approach we  
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1 clearly observed multiple transitions of PDK1 between the two conformations differing in the  
2 orientation and folding of helix  $\alpha$ B, with PS210 suppressing the population of the molecules  
3 with a more disordered helix. In Figure 3, we report the free energy projected along the  
4 variable describing the length of the helix  $\alpha$ B (end-to-end distance) and the distance between  
5 the Gly-rich loop (residues 90-94) and Asp205, which describes the kinase hinge motion. It is  
6 clear that the two conformations observed in the previous MD of the kinase with ATP that are  
7 well resolved along the  $\alpha$ B end-to-end distance variable are not observed in the presence of  
8 PS210, whose free energy surface (FES) has a single minimum with a well-ordered  $\alpha$ B helix  
9 (Fig. 3B). The second minimum of the FES of PDK1 with ATP is also compatible with a lower  
10 degree of  $\alpha$ B integrity and with our observation that partial unfolding of the C-terminal end of  
11 the helix takes place. Although the FES along the Gly-rich loop-Asp205 distance (hinge  
12 motion) shows a clear main minimum for ATP and ATP + PS210, the morphology and  
13 position of the basins suggests that the binding of PS210 restricts the kinase lobe dynamics,  
14 enforcing a more closed catalytic domain (Fig. 3B), in agreement with the crystal structure of  
15 the complex (Busschots et al., 2012). Indeed, structures with a Gly-rich loop-Asp205 distance  
16 larger than 20 Å, appear to be relatively well populated in the presence of ATP, but are rarely  
17 populated upon addition of PS210. A highly conserved salt bridge between a Lys that  
18 positions the phosphate of ATP at the active site and a Glu from the helix  $\alpha$ C (Lys111 and  
19 Glu130 in PDK1) is widely considered a hallmark of active structures of protein kinases. The  
20 MD simulations show that in the presence of ATP, the Lys111-Glu130 salt bridge is present  
21 in closed structures. In agreement with being an activator, PS210 stabilizes conformations  
22 with a closed hinge and well-formed helix  $\alpha$ B. The same conclusion can be drawn by  
23 observing the projection of the FES along the first two eigenvectors of the principal  
24 component analysis (PCA) (Fig. S4D). Different orientations of Tyr126 were observed in our  
25 new simulations for PDK1 with both ATP and PS210 (Fig. S4C and Fig. S5A). Interestingly,  
26 the same alternative conformation of Tyr126, with the OH in close proximity to the terminal  
27 phosphates of ATP, was observed independently in a new crystal form of PDK1 obtained in  
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1 complex with ATP and a phosphorylated HM polypeptide binding to the PIF-pocket (Fig. S4C  
2 and Table S1). In addition, this new crystal structure confirms the PIF-pocket as the binding  
3 site of substrate HM polypeptides (Fig. S5B). Overall, the PT-WTE simulations confirmed the  
4 site of substrate HM polypeptides (Fig. S5B). Overall, the PT-WTE simulations confirmed the  
5 differences in the behavior of ATP- and ATP-PS48/PS210-bound PDK1 previously observed  
6 by MD and allowed better characterization of the dynamic and structural shifts arising from  
7 the binding of the allosteric effector. Together, the molecular dynamics simulations were in  
8 excellent agreement with the experimental data, and predicted Tyr126 movements that were  
9 independently confirmed in successive crystallography work. Noteworthy, the simulations in  
10 the presence of ATP revealed that in the open conformation the PIF-pocket is always well  
11 formed and stable, while the closed structure presents also a destabilized  $\alpha$ B helix.  
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### 26 **Molecular mechanism of the reverse allosteric effects of PS653 and adenosine on the** 27 **PIF-pocket**

28 We next analyzed the molecular mechanism of the reverse allosteric effects of PS653 and  
29 adenosine on the PIF-pocket. The free energy projections along the same variables used  
30 before are reported in Figure 3B. PS653 produced a marked enhancement of the PDK1  
31 hinge/twist motions (Fig. 3B and Fig. S4D). This, in turn, dramatically increases the  
32 population of more open structures (distances larger than 17Å in Figure 3B). In contrast,  
33 adenosine enriched the most closed conformation (Fig. 3B), suppressing the hinge motion,  
34 similar to what was observed in the PS210 FES. While both PS210 and PS653 rigidify the  
35 PIF pocket, the most populated conformations are quite different, and the two lobes appear  
36 more twisted in the structure with PS653 (Fig. S6A,B). The additional hydrogen bonds  
37 formed by the ribose moiety of adenosine (Fig. 2C) could trigger the hinge motion and tip the  
38 balance of the conformational equilibrium towards more closed. Moreover, the changes  
39 observed in the conformational free energy landscape suggest that the stabilization of more  
40 closed structures by adenosine enhances the allosteric coupling between the two lobes and  
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1 affect the binding of PIFtide to the PIF-pocket by increasing the flexibility of the  $\alpha$ B region. In  
2 contrast, PS653 has unfavorable close contacts in the crystal structure and in the MD  
3 simulation induces the displacement of Leu88, hindering the full closure of the two lobes and  
4 breaking the catalytic spine (Kornev et al., 2006). We next compared the interaction of  
5 PDK1<sub>50-360</sub> with PS653 and adenosine using isothermal titration calorimetry (ITC). The results  
6 confirmed in both cases the existence of a single binding site for the compounds in solution,  
7 with dissociation constants in the low micromolar range (Fig. S7 and Table 1). Notably, the  
8 interaction of PDK1<sub>50-360</sub> with PS653 was less enthalpic and more entropy driven (-1.2  
9 kcal/mol;  $\Delta H/\Delta G = 15\%$ ) than the interaction with adenosine (-3.2 kcal/mol;  $\Delta H/\Delta G = 50\%$ ), in  
10 agreement with the above data showing that PS653 establishes less hydrogen bonds with  
11 residues in the enzyme active site and that it promotes the hinge movement stabilizing the  
12 open conformation of the catalytic domain. Thus, the presence of PS653 tips the balance of  
13 the conformational equilibrium towards more open structures and in turn to a specific  
14 conformation of the PIF-pocket that is less apt at binding PIFtide. This finding also suggests  
15 that the conformational flexibility of the  $\alpha$ B region might be required for proper recognition of  
16 PIFtide and HM polypeptides of other physiological substrates that dock in the PIF-pocket.  
17 Overall, the results indicated that the reverse allosteric effects that enhance the binding of  
18 PIFtide are transduced through the two lobes when the hinge motion brings them together in  
19 the closed conformation, while the decrease in binding to PIFtide is due to an induced  
20 opening of the hinge that interrupts the allosteric network.  
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45 Together, the computational results point towards a central role played by the hinge  
46 and twist motions and an allosteric communication network that extends from one lobe to the  
47 other. Numerous residues increase interactions in the closed conformation, including  
48 residues from the  $\alpha$ B and  $\alpha$ C helices, the DFG motif, the Gly-rich loop as well as residues  
49 from the large lobe (Fig. S6C). In addition, the molecular dynamics simulations identified a  
50 series of salt bridges that were formed or broken with the hinge motion, e.g., Glu130-Lys111,  
51 Asp138-Lys199, Asp138-Lys144 and Lys163-Glu215. Of these, the salt bridge formed by  
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1 Lys144 and Asp138 was only observed in the closed form but not in the open form, where it  
2 is replaced by the Lys199-Asp138 interaction (Fig. 4A). A mutagenesis study based on the  
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4 simulations confirmed a major role for Lys144. Firstly, we found that the purified Lys144Glu  
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6 and Lys144Ala mutant proteins had slightly lower specific activities (approximately 30%  
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8 decreased) than the control GST-PDK1 protein. In spite of lower basal activity, the Lys144  
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10 mutants were normally activated by an excess of PIFtide (2  $\mu$ M), reaching 93% and 89%,  
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12 respectively, of the wild type activity in the presence of PIFtide, indicating that they were  
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14 active and overall well folded, and suggesting that the mutation stabilized open-inactive  
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16 forms of the kinase. Secondly, the Lys144 mutations importantly decreased the binding of  
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18 PIFtide (Fig. 4B). Together, we validated that Lys144 indeed played an important role in the  
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20 hinge motion, stabilizing open conformations that have decreased ability to bind PIFtide. This  
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22 is in remarkable agreement with MD simulations of the Lys144Glu mutant. Starting from a  
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24 structure where the original Asp138-Lys144 salt bridge was formed, the Lys144Glu mutation  
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26 quickly leads to the formation of the alternative Lys199-Asp138, observed only in the open  
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28 structures of the wild type. Thus, the mutation actively shifts the population towards more  
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30 open structures that, as previously demonstrated, are generally less prone to bind PIFtide, as  
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32 the mutant itself. The mutated Glu144 doesn't appear to form any interaction and remains  
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34 exposed to the solvent for the remainder of the simulation, suggesting that the effect of the  
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36 mutation is not due to new interactions, but to the severing of important interactions in the  
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38 existing allosteric network. In spite of the much lower affinity for PIFtide, we could perform a  
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40 suitable interaction assay upon increasing the concentration of biotin-PIFtide in the assays  
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42 (Fig. 4D,E). Notably, the PDK1 Lys144 mutants increased the binding of PIFtide in the  
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44 presence of adenosine (Fig. 4D). In contrast to the lack of reverse allosteric effects by  
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46 adenine on wild type PDK1, adenine induced very high reverse allosteric induction of the  
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48 binding of PIFtide in the PDK1 Lys144Glu mutant (Fig. 4E). As the increased population of  
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50 open structures is generally a consequence of the higher inter-lobe dynamics (as seen for  
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52 PS653), the effect seen in the Lys144Glu PDK1 mutant could be due to adenine shifting  
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back the equilibrium towards more closed structures to maximize the interactions within the  
binding site, *de-facto* restoring the open-closed equilibrium of wild type PDK1. On the whole,  
we conclude that Lys144 plays a key role in the regulated hinge movement, which is central  
to the mechanism of allosteric coupling between the ATP-binding site and the PIF-pocket.

## 10 **Drugs in development binding at the ATP-binding site and allosterically affecting the** 11 **PIF-pocket**

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14 We further investigated whether our findings can explain observed selective in-cell inhibition  
15 of downstream signaling by some PDK1 inhibitors. UCN01 is the 7'OH derivative of  
16 staurosporine, which binds to the ATP-binding site in PDK1(Komander et al., 2003;  
17 Takahashi et al., 1989) and inhibits the phosphorylation of substrates. The recently  
18 developed GSK2334470 is a very potent ATP-competitive inhibitor of PDK1(Axten et al.,  
19 2010) that shows high selectivity for PDK1. Intriguingly, GSK2334470 very efficiently inhibits  
20 S6K phosphorylation in cells but is a relatively poor inhibitor of the phosphorylation of  
21 another PDK1 substrate, PKB/Akt (Najafov et al., 2011). The reason for this differential  
22 inhibition of PDK1 substrates has remained elusive (Rettenmaier et al., 2014). Early studies  
23 described that the PIF-pocket of PDK1 is required for the phosphorylation of S6K but not  
24 PKB/Akt (Biondi et al., 2001), and this was confirmed in knock-in cells and tissues  
25 expressing PDK1[Leu155Glu] (Arencibia et al., 2013; Bayascas, 2008; Collins et al., 2003)  
26 as well as pharmacologically using compounds binding to the PIF-pocket of PDK1, which  
27 block the phosphorylation of S6K but not PKB/Akt (Busschots et al., 2012; Rettenmaier et al.,  
28 2014). Given that S6K but not PKB/Akt requires a docking interaction of its HM to the PIF-  
29 pocket of PDK1 for its efficient phosphorylation, we hypothesized that the differential effect of  
30 GSK2334470 on S6K and PKB/Akt could be related to the identified reverse allosteric  
31 regulation effect. We performed a set of experiments with both GSK2334470 and UCN01  
32 and observed that only the GSK compound displaced the binding of PIFtide from PDK1 (low  
33 nM IC50), acting similar to PS653, while UCN01 did not affect the binding of PIFtide even at  
34 high  $\mu$ M concentrations (Fig. 5A,B). The results suggest a model in which the reverse

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allosteric effect induced by GSK2334470 enhances the inhibition of S6K phosphorylation, which requires binding to the PIF-pocket. In this manner, GSK2334470 could inhibit S6K through a dual mechanism, inhibiting the intrinsic activity of PDK1 and additionally disturbing the docking interaction by reverse allostery, whereas PKB/Akt phosphorylation would be affected by the inhibition of PDK1 intrinsic activity but not by the reverse allosteric effect.

The above findings enable the development of substrate selective inhibitors of kinases by targeting the ATP-binding site. In order to better illustrate possible uses of the described mechanism we tested the existence of such effects on the mitotic protein kinase Aurora A. The site equivalent to the PIF-pocket in Aurora A binds to a protein partner, TPX2, a microtubule associated protein which localizes the kinase in the centrosomes during mitosis. Aurora kinase also forms complexes with the oncogenes N- and C-Myc, thereby stabilizing Myc and supporting tumor growth. There is interest in the identification of drugs that bind to Aurora kinase, disturb the complex and destabilize the Myc oncogene (Brockmann et al., 2013; Gustafson et al., 2014). We have thus set up Aurora A-TPX2tide interaction assays, similar to those developed to test the interaction between PDK1 and PIFtide. Two different ATP-competitive drugs that have entered clinical trials were then tested on the reverse allosteric effect in Aurora kinase. Interestingly, while VX680 virtually did not effect the interaction (at  $\mu\text{M}$  concentrations), MLN8237 potently displaced the interaction (at low nM concentrations) (Fig. 5C,D). The finding on Aurora kinase indicates that different compounds binding to the ATP-binding site can produce very different reverse allosteric effects on TPX2, an approach that can be exploited for the destabilization of N- and C-Myc as well as for the inactivation of Aurora kinase by delocalization.

## Discussion

The last decade has seen increasing interest in the development of allosteric drugs. However, the first examples of the rational modulation of allosteric transitions of proteins by

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small compounds have only recently started to emerge (Herbert et al., 2013). We here describe small compounds that take advantage of the bi-directional nature of allosteric networks, enhancing or inhibiting the binding to a downstream signaling partner and further defined the molecular mechanism for the reverse allostery between the active site and the PIF-pocket in PDK1 (Fig. 6). This knowledge can be exploited for protein kinase drug discovery and drug development.

We not only describe compounds that make full use of the bi-directional nature of allosteric communication by binding at the ATP-binding site and affecting the PIF-pocket regulatory site, but we are also depicting the global molecular mechanism underlying the allosteric cross-regulation of the two sites. The hinge motion plays a pivotal role in allowing the allosteric communication. Orthosteric compounds (e.g. adenosine) that re-modulate the hinge motion and bring the lobes together promote the formation of an allosteric network connecting the ATP-binding site and the PIF-pocket. Various residues pertaining to the catalytic and regulatory spines (C- and R-spines) are part of the network (Kornev et al., 2006), and two salt bridges alternatively formed by Lys144 act as a switching mechanism turning the communication on or off. Adenosine by allosterically increasing the flexibility of  $\alpha$ B and  $\alpha$ C helices (Figure 4B) increases the ability of the PIF-pocket region to adapt to the binding PIFtide peptide. In stark contrast to adenosine, PS653 weakens the C-spine and favors an open-hinge conformation, breaking the allosteric communication network and locking the  $\alpha$ B and  $\alpha$ C helices lining the PIF-pocket in a conformation less favorable to the binding of the PIFtide peptide. In this manner, PS653 disfavors the binding of the prototypical hydrophobic motif polypeptide that physiologically bind to the PIF-pocket.

Our initial unbiased MD simulations clearly showed the existence of different conformations of PDK1, which fitted very well to all previous knowledge on the biochemical and structural information on the activation of PDK1 by PIFtide and small compounds binding to the PIF-pocket. With current availability of computer resources it is not possible to extend

1 such unbiased simulations up to the high  $\mu$ s to ms time scales. However, this unbiased result  
2 prompted us to explore the dynamics of PDK1 by focusing on particular areas of the protein,  
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4 by using PT-WTE, an effective enhanced-sampling algorithm. The detailed information from  
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6 biochemical studies, structural work and molecular dynamics simulations (MD and PT-WTE)  
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8 provided a clear mechanism of the bi-directional allosteric process in PDK1. We believe that  
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10 a similar approach can be effectively used to investigate the mechanisms of allosteric  
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12 regulation by occupancy of the site equivalent to the PIF-pocket in other protein kinases.  
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16 The previous characterization that GSK2334470 is a much stronger inhibitor of S6K  
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18 than of PKB/Akt could be explained by the fact that this compound does not produce a  
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20 double inhibitory effect on PKB/Akt, because this kinase does not require the HM-docking  
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22 interaction with the PIF-pocket of PDK1 for its physiological phosphorylation. Thus, at  
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24 concentrations where PDK1 is not fully inhibited by GSK2334470, its effects are similar to  
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26 compounds binding to the PIF-pocket, which selectively inhibit S6K, without affecting  
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28 PKB/Akt (Busschots et al., 2012; Rettenmaier et al., 2014). There is some precedence for  
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30 paradoxical effects observed by drugs binding at the ATP-binding site of protein kinases. A  
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32 reverse allosteric effect is probably responsible for the nucleotide pocket-induced activation  
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34 of PKC $\epsilon$  (Cameron et al., 2009) and the allosteric RNase activation of IRE1 $\alpha$  (Wang et al.,  
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36 2012). In addition, there is some evidence that a similar reverse allosteric modulation might  
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38 be involved in the “paradoxical activation” of partially inhibited B-RAF dimers (Heidorn et al.,  
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40 2010). Interestingly, the detailed characterization of the allosteric modulation by compounds  
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42 binding to the ATP-binding site could inspire the rational development of drugs with add-on  
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44 characteristics. A rich example to discuss the implications is that of Aurora kinase.  
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46 Compounds that bind to the ATP-binding site and allosterically affect the interaction with  
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48 TPX2 are expected to destabilize the oncogene Myc by destabilizing the Aurora-Myc  
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50 complexes. In addition, since TPX2 provides proper localization to Aurora A, compounds that  
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52 bind to the ATP-binding site and allosterically disrupt the interactions with TPX2, may render  
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54 molecules functionally unable to phosphorylate their specific substrates, even after the  
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1 dissociation of the compound. Similarly, it can be expected that compounds that produce  
2 desired double effects, i.e blocking the active site and, in addition, disrupting or enhancing a  
3 given allosteric interaction may be desired. We must note that the current nomenclature to  
4 describe the mode of inhibition of kinases by small compounds (type I, type II and type III  
5 inhibitors) does not appropriately describe the allosteric effects of the compounds. A more  
6 appropriate nomenclature, in use for allosteric compounds acting on receptors, requires  
7 naming the identity of the receptor, the binding site of the small molecule, the site which is  
8 being allosterically affected and the kind of modulation. In one of the examples described  
9 here, GSK2334470 is a type I inhibitor but is also a negative allosteric modulator binding to  
10 the ATP-binding site and displacing the binding of PIFtide from the PIF-pocket of PDK1.  
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23 Here, we also provide the crystal structure of PDK1 in complex with the  
24 phosphorylated HM of Akt/PKB. The equivalent docking interaction is necessary for the  
25 phosphorylation and activation of most PDK1 substrates, including S6K and SGK. However,  
26 the PIF-pocket docking interaction is not considered to be required for the phosphorylation of  
27 Akt/PKB (Biondi et al., 2001; Busschots et al., 2012; Collins et al., 2003; Rettenmaier et al.,  
28 2014). Our structure suggests that the docking interaction still takes place for Akt/PKB, even  
29 if it is not the determinant of the phosphorylation of this kinase. While writing the current  
30 manuscript, the crystal structure of PDK1 in complex with a short version of PIFtide was  
31 published (Rettenmaier et al., 2014). From our results, it appears that PDK1 might require a  
32 limited increased flexibility of the PIF-pocket to sample the appropriate conformation for  
33 PIFtide binding. According to this scenario, the disorder of helix  $\alpha$ C previously observed in  
34 solution (Hindie et al., 2009) and the disorder of helix  $\alpha$ B predicted here by the molecular  
35 dynamics simulations might provide a mechanism to facilitate the binding of different  
36 substrate HM polypeptides. This entropy-driven binding mechanism, also observed in ternary  
37 complexes of PKA (Masterson et al., 2011), may be an essential component of the ability of  
38 PDK1 to act as a “conformational sensor” (Biondi, 2004).  
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2 The initial crystal structures of PKA already depicted the versatility of the kinase  
3 domain revealing open, intermediate and closed conformations of the ATP-binding site  
4 (Johnson et al., 2001). The hinge interlobe motion, was afterwards described by molecular  
5 dynamics simulations of PKA (Masterson et al., 2011) and other kinases (Dolker et al.,  
6 2014). In PKA, it was revealed that the rate of hinge motion of PKA catalytic domain ( $20 \text{ s}^{-1}$ )  
7 correlated with the turnover of the enzyme (Kim et al., 2015; Srivastava et al., 2014). The  
8 turnover of PDK1 phosphorylation of different substrates, however, is approximately 1000  
9 times slower ( $0.6\text{-}1 \text{ min}^{-1}$ ) (Biondi et al., 2001), therefore indicating that most opening and  
10 closing motions of the PDK1 kinase domain are non-productive, not leading to the  
11 phosphorylation of substrates.  
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23 It is worth noting that a very important cellular molecule such as adenosine produces  
24 reverse allosteric effects on PDK1 *in vitro*. Adenosine is not only a metabolite from the  
25 synthesis of ATP, the substrate of PDK1, but it is also a signaling molecule on its own (Borea  
26 et al., 2016). Our results suggest that the increase in adenosine could lead to the  
27 engagement of PDK1 in complexes with S6K, subtly regulating the identity of substrates of  
28 PDK1 to become phosphorylated. It is tempting to speculate that other protein kinases could  
29 as well respond to the levels of ATP intermediates or other small physiological nucleotide  
30 molecules. This mechanism could indeed regulate cellular signaling by affecting the  
31 interactions of proteins with regulatory domains or regulatory subunits, formation of protein  
32 complexes, modulation of localization or phosphorylation by upstream kinases, etc  
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46 Previous data indicates that the full bi-directional allosteric regulation between the  
47 active site and the PIF-pocket regulatory is functional in the regulation of AGC kinases,  
48 perhaps best exemplified in PRK2, where the a pseudosubstrate inhibitory polypeptide,  
49 PLKtide binding to the active site, and PIFtide binding to the PIF-pocket, are allosterically  
50 mutually excluded and contribute to PRKs complex mechanism of regulation (Bauer et al.,  
51 2012).  
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Although the mechanisms of allostery have been investigated for the last 50 years, it is still not possible to rationally design small molecules to modulate allosteric cellular switches at will. The present work provides a deep understanding of the bi-directional allosteric coupling mechanism and suggests ways to exploit it. The use of molecular dynamics simulations as described above enables the *in silico* testing of molecules before synthesis and provides a technology to guide the design of variant compounds that produce a desired reverse allosteric effect. We expect that the results from this work will inspire the development of drugs with reverse allosteric effects on other protein kinases and, more generally, on other signaling proteins.

### Significance

Allostery is a fundamental and widespread regulation mechanism by which proteins transfer information between remote sites and functional sites in response to different stimuli. It is of interest to modulate allosteric processes for treatment of diseases and for future design of regulated molecular machines. Our proposed PDK1 model provides information on the conformational transitions of the catalytic domain and depicts for the first time, in atomistic details, the bidirectional allosteric communication between the ATP-binding site and the regulatory PIF-pocket, where compounds binding to the ATP-binding site can allosterically enhance or inhibit the interaction with a cellular partner. More generally, drugs directed to the ATP-binding site that have equal ability to inhibit a given kinase, may have very different effects on the formation of protein complexes and cell signaling, different on-target side effects and overall different success in the treatment of patients. The PDK1/AGC kinase allosteric model system provides a rich example of the potential of small molecules to bi-directionally modulate the conformation of a protein kinase. By providing a validated computer simulation it will now be possible to use this tool to design allosteric drugs with a given allosteric effect. In turn, we expect that the perspective presented here will benefit drug development to members of the protein kinase family by providing a computer platform from where to rationally design compounds directed to the PIF-pocket or ATP-binding site and

1 producing desired secondary allosteric effects. Similarly, the above approach could also be  
2 applied more generally for drug discovery projects to other allosteric signaling proteins  
3 whose function could be modulated by molecules designed to produce desired  
4 conformational changes.  
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## 10 11 **Experimental Procedures**

### 12 13 **Materials**

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15 The polypeptide substrate of protein kinase PDK1 was T308tide (KTFCGTPEYLAPEVRR; >  
16 75% purity). Other polypeptides used were PIFtide (REPRILSEEEQEMFRDFDYIADWS),  
17 biotin-PIFtide (biotin-REPRILSEEEQEMFRDFDYIADWS) and biotin-TPX2tide (Biotin-  
18 MSQVKSSYSYDAPSDFINFSSLDDEGDTQNIDSWFEEKANLEN-NH<sub>2</sub>). Adenine and  
19 adenosine (≥ 99%) were from Sigma-Aldrich.  
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### 29 30 **Crystal structures**

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32 To obtain the crystal structure of PDK1 in complex with PS653, PSE10, adenine, and  
33 adenosine, PDK1 was expressed, purified, concentrated, crystallized and soaked with  
34 compounds as previously described (Hindie et al., 2009). The crystal structure of PDK1 in  
35 the new crystal packing was obtained in a screening for new crystallization conditions in the  
36 presence of HM-polypeptides. PDK1<sub>50-359</sub> [Y288G,Q292A] was co-crystallized in the  
37 presence of the phosphorylated peptide KGAGGGGFPQFS(P)YSA (underlined residues are  
38 conserved in the HM phosphorylation site of Akt/PKBs, SGKs, S6Ks, and PKCs). The  
39 coordinates of the structures have been deposited in the Protein Data Bank under accession  
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### 53 54 **PDK1 expression, purification, and in vitro activity test**

55 His-tagged PDK1<sub>1-556</sub> and His- tagged PDK1<sub>50-359</sub> employed in activity assays and in the  
56 alphascreen interaction assays were expressed in insect cells and purified through Ni-NTA  
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1 and gel filtration chromatography, as described (Busschots et al., 2012; Engel et al., 2006).  
2 His-tagged human Aurora A<sub>1-403</sub> was recombinantly expressed in bacteria from pET28  
3 plasmid and purified as described for PDK1 above. The GST-fusion proteins were obtained  
4 from HEK293 cells after transient transfection of the corresponding pEBG2T plasmids and  
5 purified as described (Engel et al., 2006). The detailed conditions for the activity assay are  
6 described in the Supplemental Information.  
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### 13 **Alphascreen interaction-displacement assays**

14 The AlphaScreen assay was performed according to the standard protocol from the  
15 manufacturer (Perkin Elmer). The set-up of the interaction-displacement assay was  
16 previously described (Busschots et al., 2012). In short, the assay was performed in a final  
17 volume of 25 µl in white 384-well microtiter plates (Greiner) with His-PDK1<sub>50-359</sub> [Tyr288Gly;  
18 Gln292Ala] (25 nM) and Biotin-PIFtide (25 nM) in a buffer containing 50 mM Tris-HCl (pH  
19 7.4), 100 mM NaCl, 1 mM dithiothreitol, 0.01% (v/v) Tween-20 and 0.1% (w/v) BSA, followed  
20 by the addition of 5 µl of beads (nickel chelate-coated acceptor beads and streptavidin-  
21 coated donor beads; 20 µg/ml final concentrations). The set-up for Aurora A-TPX2tide  
22 interaction assay was the same, except that His-Aurora A (5 nM) and biotin-TPX2tide (10  
23 nM) were employed.  
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### 40 **Molecular Dynamics**

41 We performed Molecular Dynamics (MD) simulations using the GROMACS 4.6 package  
42 (Hess et al., 2008) and the Amber99SB\*-ILDN force field (Best and Hummer, 2009; Lindorff-  
43 Larsen et al., 2010). The system was minimized with 10000 steps of conjugated gradient and  
44 equilibrated in the NPT ensemble for 10 ns. A production run of 400 ns was then performed  
45 in the NVT ensemble, with a time step of 2 fs. Neighbor searching was performed every 5  
46 steps. The PME algorithm was used for electrostatic interactions with a cut-off of 1.2 nm,  
47 while a single cut-off of 1.2 nm was used for Van der Waals interactions. Temperature  
48 coupling was done with the V-rescale algorithm (Bussi et al., 2007). The parallel-tempering  
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1 simulations were performed using Gromacs 4.6(Hess et al., 2008) combined with the  
2 PLUMED 2.1 plug-in (Tribello et al., 2014). We performed PT with 5 replicas in the 300K-  
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4 400K range using the Well Tempered Ensemble (Bonomi and Parrinello, 2010).  
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### 7 **Isothermal titration calorimetry**

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9 Calorimetric titrations were performed using the MicroCal iTC200 instrument (GE Healthcare  
10 Life Sciences) as previously described (Hindie et al., 2009) with the modifications indicated in  
11 the Supplemental Experimental Procedures. For the titrations of PDK1 with adenosine, the  
12 protein and the compound were prepared in 50 mM Tris-HCl (pH 7.4), 200 mM NaCl, 1 mM  
13 DTT and 1% v/v DMSO. For the titration of PDK1 with PS653 the protein and the compound  
14 were prepared in 50 mM Tris-HCl (pH 7.4), 200 mM NaCl, 1 mM DTT and 5% v/v DMSO.  
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16 Errors on the thermodynamic parameter values in Table 1 are non linear least square fitting  
17 errors of the experimental binding isotherms using the Levenberg-Markardt iteration method  
18 (Freire et al., 2009).  
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### 30 **Small molecules**

31 Compounds **PS653** and **PSE10** were commercial available from Maybridge. The commercial  
32 compounds were further analyzed by <sup>1</sup>H-NMR, <sup>13</sup>C-NMR and ESI-MS (see Supplemental  
33 Experimental Procedures).  
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### Author contributions

JOS obtained and analyzed most of the crystallography data. VH solved initial PDK1 crystals. GS and FLG designed the computational analysis. GS did the MD simulations. KB set up the screening assay and MNL did biophysical assays. DO provided medicinal chemistry support. Screening and biochemical characterizations were performed by ES, KB, AKH and SN under the supervision of RMB. SZ provided advice. PMA supervised VH and analyzed data. RMB supervised the whole research project. The manuscript was written by RMB and FLG supported by JOS, GS and PMA.

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## Figure Legends

1  
2 **Fig. 1.** The kinase domain, the PIF-pocket binding to PIFtide, and the identification of small  
3 molecules that displace PIFtide from the PIF-pocket. **(A-B)** Crystal structure of the catalytic  
4 domain of PDK1 in complex with ATP (yellow carbon atoms) binding at the active site and  
5 PS210 (orange carbon atoms) binding at the PIF-pocket allosteric site (PDB ID: 4AW1). **(C)**  
6 Schematic representation of the alphascreen interaction assay employed for the screening of  
7 the library of small molecules. **(D)** Effect of PSE10 and PS653 on the interaction between  
8 His-PDK1 50-360 and biotin-PIFtide. **(E)** Structures of PSE10 and PS653 in comparison to  
9 previously described activators of PDK1, PS48 and PS210. **(F)** Schematic representation of  
10 the PDK1 *in vitro* activity assay, using the polypeptide T308tide as substrate of PDK1. **(G)**  
11 Effect of PSE10 and PS653 on the *in vitro* activity of His-PDK1 50-556; the inhibition of other  
12 PDK1 constructs is presented in figure S3. **(H-J)** Effect of PSE10, PS653 and PIFtide on the  
13 activity of GST-PDK1 1-556 and the PIF-pocket mutants Leu155Glu and Leu127Val. See  
14 also Figure S1-S2.

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33 **Fig. 2.** PS653, adenine and adenosine bind to the ATP-binding site but produce distinct  
34 allosteric effects at the PIF-pocket. **(A-C)** Crystal structures of PDK1 in complex with PS653  
35 **(A)**, adenine **(B)** and adenosine **(C)**. Only residues in direct contact with the compounds are  
36 shown as sticks. Hydrogen bonds are visualized as gray dotted lines. Close contacts of  
37 PS653 mentioned in the main text are indicated as red dotted lines.  $|2F_o - F_c|$  electron density  
38 of the compounds is shown in blue and contoured at  $1\sigma$ . **(D)** Effect of adenine and adenosine  
39 on the interaction between PDK1 and PIFtide. The interaction between His-PDK1<sub>50-359</sub> and  
40 biotin-PIFtide was measured using the alphascreen assay depicted in Figure 1C and the  
41 effect of adenine and adenosine quantified. See also Figure 1C and Table S1.

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55 **Fig. 3.** Free energy calculation of PDK1 with compounds. **(A)** Schematic representation of  
56 parameters measured. Free energy surfaces as projected along the identified  $\alpha B$  parameter  
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1 (end-to-end distance) and two distances describing the opening and rotation of the small  
2 lobe with respect to the large lobe. **(B)** Free energy calculation of PDK1 with ATP, PS210,  
3 PS653 and adenosine. PS653 clearly enhances the hinge motion and lobe rotation, while  
4 adenosine has the opposite effect, similar to what was observed for PS210. See also Figure  
5 S4; Figure S5.  
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13 **Fig. 4.** The reverse allosteric effect is mediated by Lys144 and is differentially affected by  
14 ATP-competitive PDK1 inhibitors. **(A)** Asp138 can form salt bridges with either Lys144 or  
15 Lys199. These residues are shown as sticks in a cartoon representation of a PDK1 crystal  
16 structure (PDB ID 3HRC). Asp138, Lys144 and Lys199 are shown. **(B-G)** The ability of GST-  
17 PDK1<sub>1-359</sub> wt and Lys144 mutants to bind biotin-PIFtide was studied in the alphascreen  
18 interaction assay. **(B)** Mutation of Lys144 to Ala or Glu affects the binding to PIFtide. **(C-E)**  
19 Effect of PS653 on the interaction between GST-PDK1proteins and biotin-PIFtide. See also  
20 Figure S6.  
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33 **Fig. 5.** Negative allosteric modulation of the binding to the PIF-pocket by drugs under  
34 development to the ATP-binding site of PDK1 and Aurora kinase. **(A-B)** Effect of UCN01 and  
35 GSK2334470 ATP-competitive inhibitors of PDK1 on the interaction of His-PDK1<sub>50-359</sub> and  
36 biotin-PIFtide. **(C-D)** Effect of VX680 and MLN8237 ATP-competitive inhibitors of Aurora  
37 kinase on the interaction of His-Aurora A and biotin-TPX2tide.  
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46 **Fig. 6.** Schematic representation of the bi-directional allosteric modulation of AGC kinases by  
47 small compounds. **(A)** Representation of the hinge motion in equilibrium between open and  
48 closed structures of the protein kinase catalytic domain, showing the orthosteric ATP-binding  
49 site (active site), the HM polypeptide PIFtide and the PIF-pocket (allosteric site). The end-to-  
50 end distance variations of the helix  $\alpha$ B are represented by the length of helices. The key  
51 residue Lys144 is represented as a red stick. **(B)** Representation of the reverse allosteric  
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1 mechanism by which PS653 and Adenosine (Ado) affect the conformation of the PIF-pocket  
2 and the binding of PIFtide. PS653 binds to the ATP-binding site, stabilizes open  
3 conformations and rigidifies the PIF-pocket, resulting in the inhibition of the binding of  
4 PIFtide. Adenosine binds to the ATP-binding site, favors the dynamic closure of the structure  
5 and disorders the PIF-pocket in the closed conformation, resulting in the cooperative binding  
6 of PIFtide. Lys144, Asp138 and Lys199 (shown as sticks) are represented in alternative salt  
7 bridge interactions formed in the closed and open conformations of PDK1.  
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**Table 1.** Thermodynamic parameters of PDK1 interaction with low-molecular-weight compounds obtained by ITC. See also Figure S7.

Compound	Temperature (K)	N	$K_a$ ( $M^{-1}$ )	$K_d$ ( $\mu M$ )	$\Delta H$ (kcal mol <sup>-1</sup> )	$\Delta S$ (cal mol <sup>-1</sup> deg <sup>-1</sup> )	T $\Delta S$ (kcal mol <sup>-1</sup> )
Adenosine	298	1.2 ± 0.2	5 10 <sup>4</sup> ± 1 10 <sup>4</sup>	20	-3.2 ± 0.7	10.7	3.2
PS653	310	0.87 ± 0.02	8 10 <sup>5</sup> ± 2 10 <sup>5</sup>	1.25	-1.20 ± 0.06	23.1	7.2

N is the stoichiometry of binding.



Figure 1

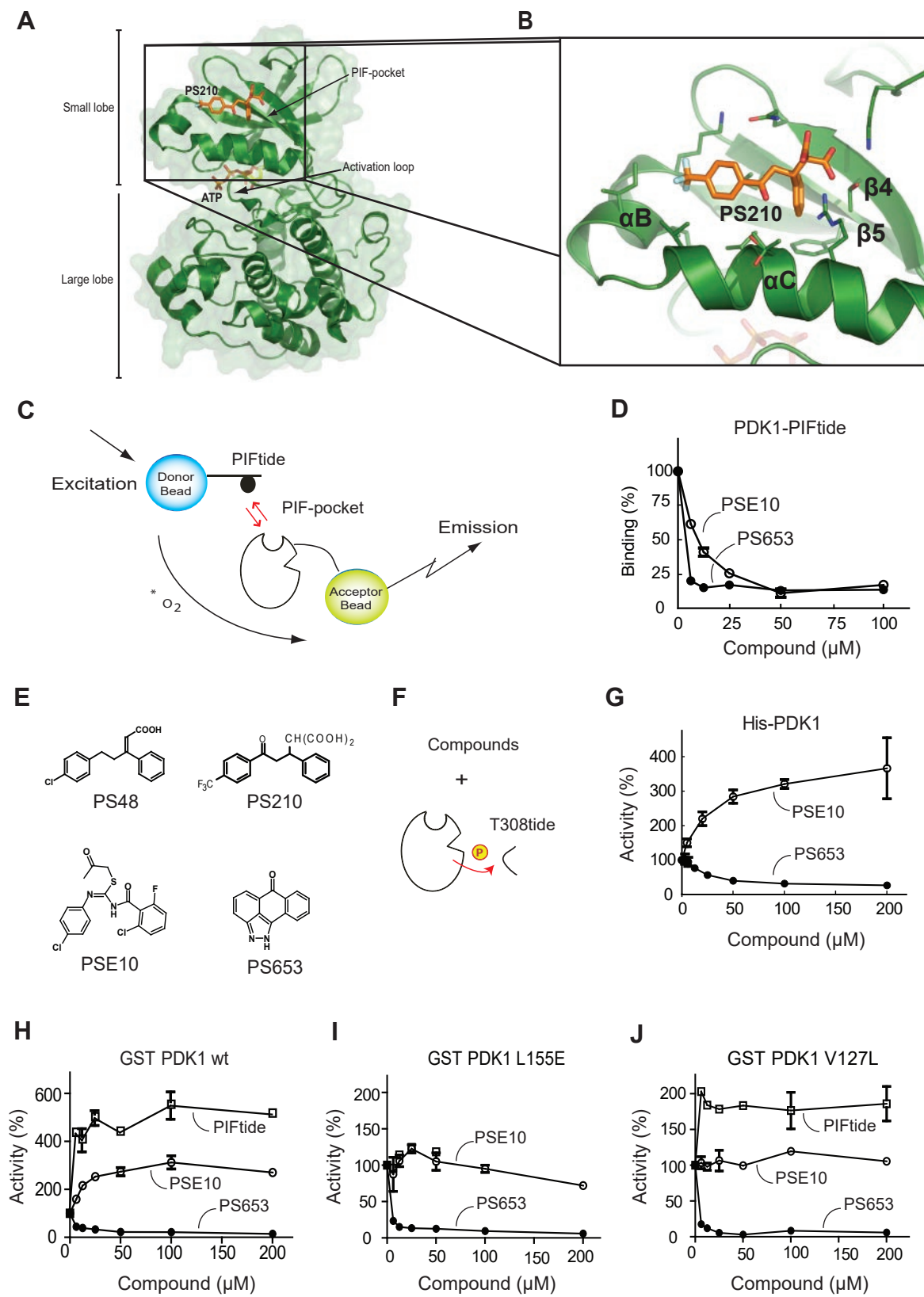


Figure 2

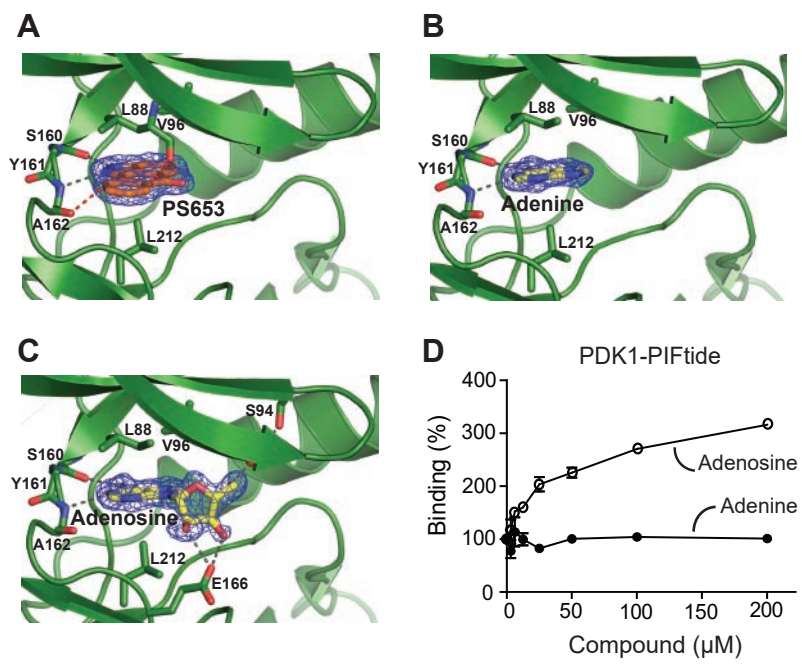


Figure 3

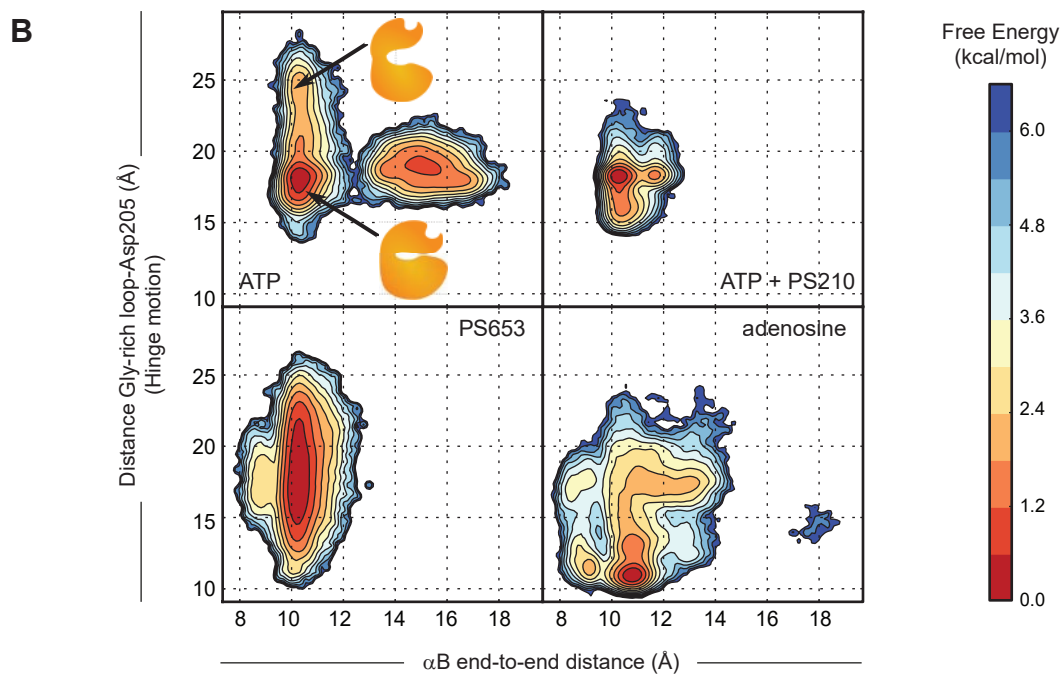
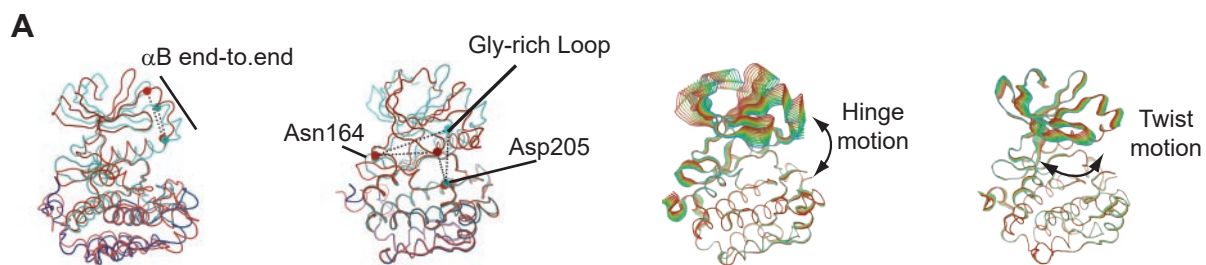


Figure 4

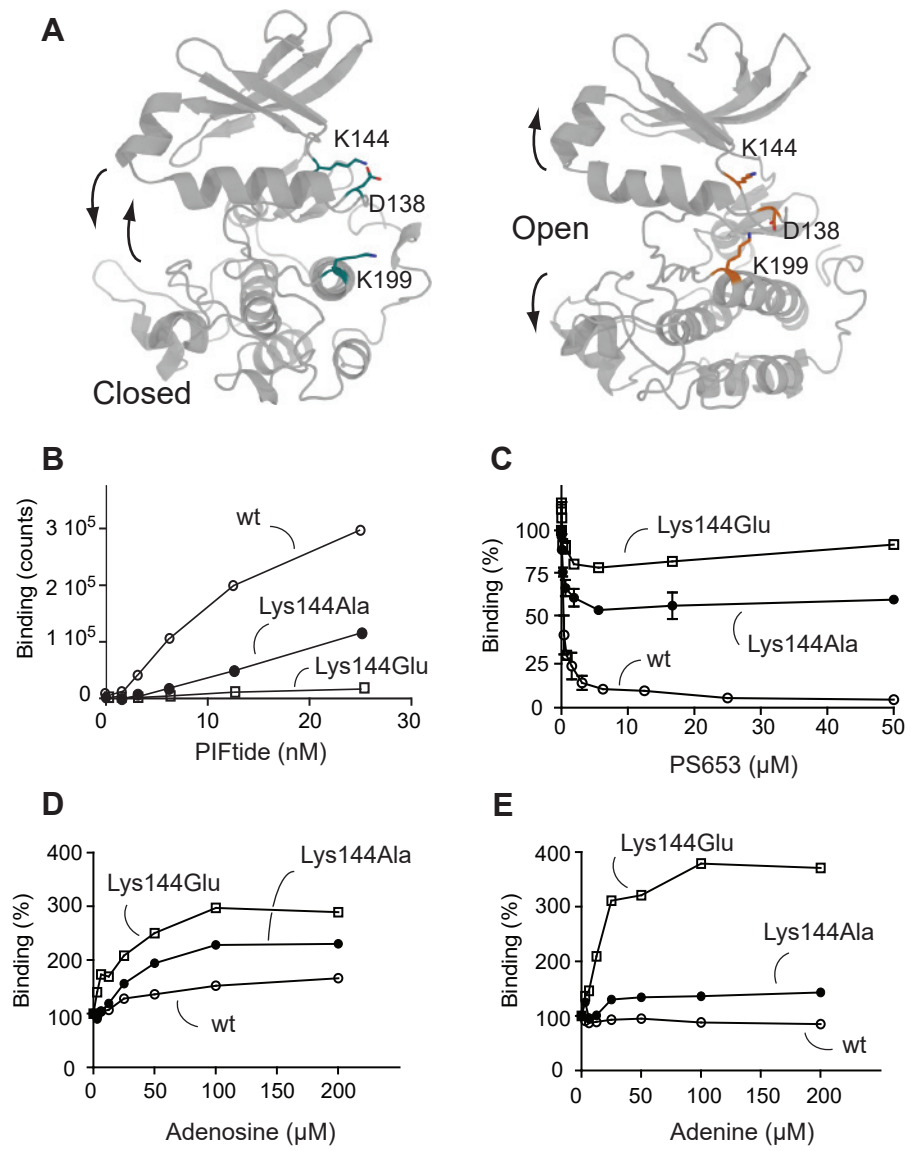


Figure 5

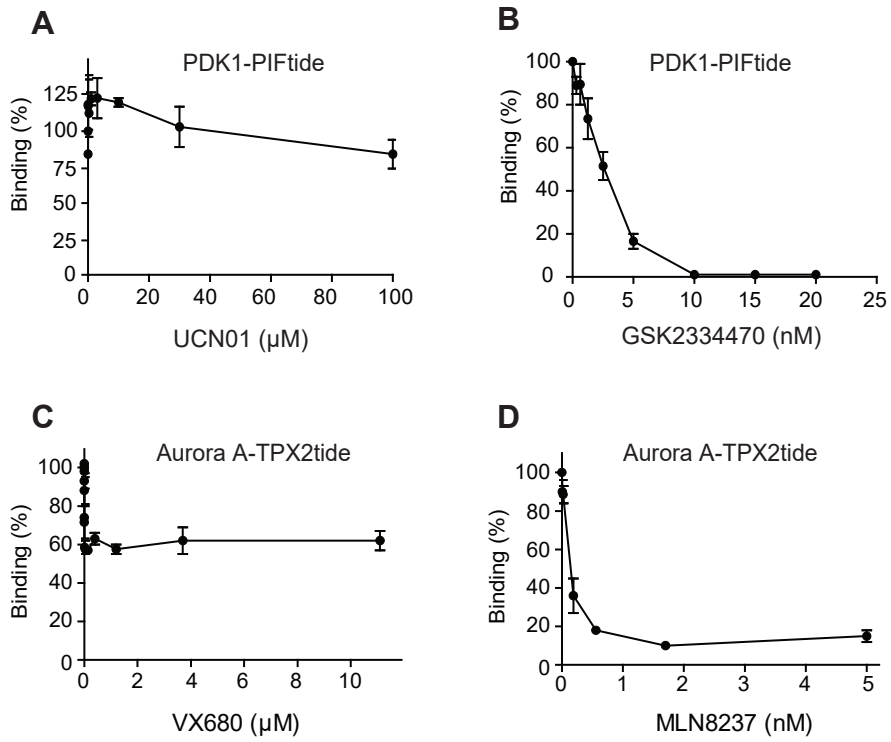


Figure 6

