Rational Redesign of a Functional Protein Kinase-Substrate Interaction

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Supporting Information

ABSTRACT: Eukaryotic protein kinases typically phosphorylate substrates in the context of specific sequence motifs, contributing to specificity essential for accurate signal transmission. Protein kinases recognize their target sequences through complementary interactions within the active site cleft. As a step toward the construction of orthogonal kinase signaling systems, we have re-engineered the protein kinase Pim1 to alter its phosphorylation consensus sequence. Residues in the Pim1 catalytic domain interacting directly with a critical arginine residue in the substrate were substituted to produce a kinase mutant that instead accommodates a hydrophobic residue. We then introduced a compensating mutation into a Pim1 substrate, the pro-apoptotic protein BAD, to reconstitute phosphorylation both in vitro and in living cells. Coexpression of the redesigned kinase with its substrate in cells protected them from apoptosis. Such orthogonal kinase–substrate pairs provide tools to probe the functional consequences of specific phosphorylation events in living cells and to design synthetic signaling pathways.

Protein phosphorylation, the most common reversible post-translational modification, is used universally as a regulatory mechanism in cellular signaling networks. Unraveling the molecular mechanisms underlying cellular regulation is dependent on our ability to experimentally control protein phosphorylation in living cells. For example, a phosphorylated residue is generally mutated to alanine to examine the consequence of preventing phosphorylation of a specific substrate protein. In addition, more elaborate synthetic signaling systems, engineered by manipulating interactions among component molecules1–5 or by introduction of artificial phosphorylation sites,6,7 can reveal basic mechanisms of signaling. Despite such impressive control over cellular signaling pathways, our capacity to simply direct phosphorylation to specific sites on proteins in living cells is remarkably limited. Phosphorylation is often simulated by introducing negatively charged Asp or Glu residues into proteins, but such mutations are necessarily constitutive and frequently fail to substitute for true phosphorylation.6 Here, we demonstrate how a protein kinase can be re-engineered by mutating its catalytic domain to phosphorylate a specific substrate in living cells. Thus far, engineering of protein kinase catalytic domains has mainly focused on the nucleotide binding site. Substitution of key residues has produced “analog-sensitive” kinase mutants that can accept modified ATP analogs and inhibitors, facilitating specific inhibition of the kinase and identification of its direct downstream substrates.7,8 By contrast, re-engineering the peptide binding specificity of kinases has been limited to a few cases primarily aimed at validating structural and evolutionary models for kinase–substrate interactions.8–14 Though not yet realized, protein kinase mutants with altered phosphorylation site specificity could potentially offer powerful tools for probing and controlling signaling pathways within living cells. For example, when expressed alone in cells, such mutants are expected to be nonfunctional, as they would not recognize and phosphorylate their endogenous substrates. However, introduction of compensating mutations into a known substrate would restore its ability to be phosphorylated by the mutant kinase. Such an orthogonal kinase–substrate pair would then allow one to directly probe the functional consequences of phosphorylation at a single site. A conceptually similar approach employing orthogonal G protein coupled receptor (GPCR)–ligand pairs (termed receptors activated by solely synthetic ligands, or RASSLs) has been used extensively for functional elucidation of GPCR signaling pathways in vivo.15

Altering the target sequence of a protein-modifying enzyme is particularly challenging, as it involves both engineering an enzyme and redesigning a protein interaction interface. We chose to use a structure-based design approach, taking advantage of available X-ray crystal structures of kinase–substrate complexes, as well as insight into the substrate preferences of many protein kinases, collected using peptide library screening methods.14,16–23 We focused our initial efforts on the serine-threonine kinase Pim1, which promotes cell growth and survival and has been implicated as an oncogenic driver in leukemia, breast cancer, and prostate cancer.24 Pim1 is thought to contribute to cell survival in part through phosphorylation of the pro-apoptotic protein Bcl2-related

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antagonist of cell death (BAD), causing the protein to be inactivated through sequestration by 14−3−3 proteins. The primary Pim1 phosphorylation site on BAD (Ser112), as well as most other known substrates of the kinase, conforms to its preferred R-x-R-x-x-S/T consensus sequence that has been established through analysis of peptide substrates. X-ray crystallography of Pim1 in complex with a consensus peptide substrate revealed four residues (Thr134, Asp170, Asp234, and Asp239) in the peptide-binding cleft in direct contact with the guanidino headgroup of the Arg residue located at the −5 position in the substrate (Figure 1a). Likewise, in the X-ray crystal structure of the MARK2, a related protein kinase selective for aliphatic hydrophobic residues at the −5 position, the corresponding Leu residue of a bound pseudosubstrate inhibitor interacted with a structurally analogous site. We therefore focused our efforts to re-engineer Pim1 specificity on mutation of the four residues within this pocket. In previous studies, we have found through peptide library screening that many kinases related to Pim1 select hydrophobic, rather than basic, residues at the −5 position. Sequence alignment of these kinases with Pim1 revealed a correlation between selectivity at the −5 position and the identity of residues found at positions analogous to the Pim1 −5 interaction pocket (Figure 1b), suggesting specific residues to introduce into the pocket to change the substrate specificity of Pim1.

To determine whether we could indeed re-engineer Pim1 to phosphorylate a new consensus sequence, we generated a series of single, double, triple, and quadruple mutants in which residues found at the analogous positions in other kinases were introduced into the −5 interaction pocket of Pim1. These mutants involved either direct grafting of residues found in a single kinase onto Pim1 or construction of a consensus kinase including residues found frequently at each of the four positions in hydrophobic-selecting kinases. In total, 12 mutant kinases were expressed, purified, and subjected to peptide library screening to assess changes in substrate specificity (Figure 2a; full peptide array analysis is shown in Figure S1). We found that mutation of the three Asp residues in the −5 pocket was sufficient to convert Pim1 from selecting primarily basic to hydrophobic-selecting kinases at the −5 position but that the quadruple mutants appeared to be the most selective in general. To quantitatively assess substrate specificity and overall level of activity, we assayed each quadruple mutant using a pair of WT and quadruple mutants on peptide substrates assessed by radiolabel kinase assay. LPVT, Pim1-T134L/D170P/D234V/D239T; LPY, Pim1-T134L/D170P/D234I/D239Y; VPIY, Pim1-T134V/D170P/D234I/D239Y; LPTY, Pim1-T134L/D170P/D234T/D239Y. Error bars indicate standard deviation (n = 3).

Figure 1. Residues in the −5 interaction pocket of Pim1 and related kinases. (A) The crystal structure of Pim1 in complex with a peptide substrate, showing the four residues (Thr134, Asp170, Asp234, and Asp239) that make direct contact with an Arg residue at the −5 position. (B) Residues found at the analogous positions in related kinases that select basic (top, shaded blue) or hydrophobic (bottom, shaded gray) residues at the −5 position on the basis of published peptide array analyses. The crystal structure of the MARK2, a related protein kinase selective for aliphatic hydrophobic residues at the −5 position, the corresponding Leu residue of a bound pseudosubstrate inhibitor interacted with a structurally analogous site. We therefore focused our efforts to re-engineer Pim1 specificity on mutation of the four residues within this pocket. In previous studies, we have found through peptide library screening that many kinases related to Pim1 select hydrophobic, rather than basic, residues at the −5 position. Sequence alignment of these kinases with Pim1 revealed a correlation between selectivity at the −5 position and the identity of residues found at positions analogous to the Pim1 −5 interaction pocket (Figure 1b), suggesting specific residues to introduce into the pocket to change the substrate specificity of Pim1.

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We next examined whether Pim1 mutation affected phosphorylation of the protein substrate BAD. As reported, WT Pim1 robustly phosphorylated recombinant BAD at Ser112 in vitro, while none of the Pim1 quadruple mutants detectably phosphorylated BAD at that site (Figure 3a). By contrast, a BAD-R107L mutant, in which the −5 Arg residue was replaced with Leu, was not a substrate of WT Pim1. However, Pim1-LPVT and, to a lesser extent, the other Pim1 mutants did phosphorylate BAD-R107L, albeit at a much lower rate compared to WT Pim1 phosphorylation of WT BAD. Introduction of an additional point mutation at the +1 position in BAD (Y113G) to more closely match the sequence preference of Pim1 as determined from peptide library screening (see Figure S1) greatly restored phosphorylation by Pim1-LPVT without providing substantial phosphorylation by WT Pim1 (Figure 3b). These results indicated that by
introducing compensating mutations into BAD, we could reconstitute its phosphorylation by mutant Pim1.

To determine whether this redesigned kinase-substrate interaction was functional, we first examined phosphorylation of BAD by Pim1 in cultured cells, focusing on the Pim1-LPVT mutant, which was most active in vitro (Figure 3c). As previously reported for other cell types,25,26 coexpression of WT Pim1 with WT BAD in HEK293 cells increased the level of Ser112 phosphorylation as assessed by reactivity with a phospho-specific antibody. As anticipated, Pim1-LPVT did not phosphorylate WT BAD at Ser112 in cultured cells. Pim1-LPVT induced phosphorylation of BAD-R107L in cells to approximately 50% of the level of phosphorylation of WT BAD by WT Pim1, in keeping with its reduced phosphorylation efficiency in vitro. Though mutation of Tyr113 abolished reactivity with the phospho-specific antibody, we could assess phosphorylation of BAD-R107LY113G in cells by electrophoretic mobility shift. Both BAD-R107L and R107LY113G underwent a Pim1-LPVT induced mobility shift that was completely eliminated by S112A mutation (Figure 3c and data not shown). This analysis indicated that Pim1-LPVT phosphorylated BAD-R107LY113G to high stoichiometry in cells, though some phosphorylation by WT Pim1 was also evident. These results indicate that our redesigned kinase-substrate system can be used to direct site-specific phosphorylation in living cells.

We next examined the effect of BAD phosphorylation on Pim1-mediated cell survival, using combinations of WT and mutant kinase and substrate (Figure 3d). BAD overexpression is reported to sensitize multiple cell types to apoptotic stimuli in a manner that can be inhibited by coexpression of kinases that phosphorylate BAD.27 We found that expression of WT BAD, the nonphosphorylatable mutant BAD-S112A, or the −5 substituted BAD-R107L and BAD-R107LY113G mutants each induced similar levels of cell death when overexpressed in COS7 cells subjected to serum withdrawal (Figure 3d and data not shown). As expected, cell death induced by WT BAD expression was largely reversed by coexpression with WT Pim1. In keeping with its inability to phosphorylate BAD-S112A or BAD-R107L, WT Pim1 was incapable of protecting cells from expression of these BAD mutants. Pim1-LPVT, by contrast, was unable to block cell death induced by WT BAD, yet rescued cells from BAD-R107L expression, correlating with its ability to substantially phosphorylate mutant BAD in cells. Unexpectedly, Pim1-LPVT did not block cell death induced by BAD-R107LY113G expression (data not shown). Pim1 inactivation of BAD may depend on Tyr113, which could be required for example for optimal binding of phosphorylated BAD to 14−3−3 proteins. Nonetheless, the ability of Pim1-LPVT to inhibit cell death specifically induced by BAD-R107L indicates that a redesigned Pim1-BAD kinase-substrate pair can be biologically functional in living cells.
In summary, we have successfully directed a functional phosphorylation event to mutant protein by means of a re-engineered protein kinase. Such an approach may find utility in elucidating the cellular consequences of individual phosphorylation events. We also note that the ability of a compensating mutation in the substrate to restore phosphorylation by the mutant provides unambiguous confirmation that a kinase-substrate interaction is direct in intact cells. Theoretically, this approach is applicable to any kinase with well-defined substrate specificity. In the example described here, we have changed the target sequence of the kinase to match that of other known kinases, which could potentially lead to “off-target” phosphorylation when the mutant kinase is expressed in cells. The use of genetic selection methods, which have been successfully applied to proteases and other enzymes, could be used in the future to confer non-native selectivity onto kinases to achieve more complete orthogonality in synthetic kinase signaling systems.

**METHODS**

**Vector Construction and Mutagenesis.** The mammalian expression vector for N-terminally GFP-tagged Pim1 was generated by PCR amplification of the full length human coding sequence and insertion into the EcoR I and Xh o sites of the vector pEGFP-C2 (Clontech). The inducible GFP-Pim1 expression vector was generated by first shuttling the entire GFP-Pim1 coding sequence into the Gateway entry vector pDONR221 by BP recombination, followed by LR recombination into the lentiviral vector pINDUCER20. The bacterial expression to produce GST-BAD-His 6 was produced by PCR amplification of the mouse BAD coding sequence with a 3’ primer encoding a His 6 tag sequence at the C-terminus and inserting into pGEX-4T2. Pim1 and BAD point mutants were generated via successive rounds of site-directed mutagenesis using the QuikChange protocol (Stratagene) with primers listed in the Supporting Information. All mutant constructs were verified by sequencing the entire open reading frame.

**In Vitro Kinase Assays.** The His 6 -tagged Pim1 catalytic domain 27 was expressed in E. coli. Detailed procedures for protein expression and purification, peptide synthesis, peptide library screening, and peptide kinase assay conditions are described in the Supporting Information. BAD kinase reactions contained 5 or 20 nM kinase and 2 μM purified BAD in kinase assay buffer. Reactions were started by the addition of ATP (to 100 μM, with or without 0.25 μCi/μL [γ-32P]ATP). Cold reactions were quenched after 5 min incubation at 30 °C by adding 4X SDS-PAGE loading buffer and heating to 95 °C for 5 min. Samples were fractionated by SDS-PAGE followed by transfer to the PVDF membrane. Blots were probed with either the α-phosphoSer112 antibody (Cell Signaling Technology #9291) or mouse α-penta-His antibody (Qiagen #34660) followed by the appropriate horseradish peroxidase-linked secondary antibody, and developed by enhanced chemiluminescence. Hot reactions were quenched in the same way after 20 min incubation at 30 °C and fractionated by SDS-PAGE. Gels were dried and radiolabel incorporation detected by phosphor imaging.

**Analysis of BAD Phosphorylation in Cultured Cells.** HEK293 cells in six-well plates were co-transfected with pEGF-Pim1 (0.25 μg for WT Pim1 or 0.5 μg for mutant Pim1 per well, adjusted to achieve equal levels of protein expression), pEBG-BAD (Cell Signaling Technology, 3.5 μg WT or mutant per well), and empty control (to bring the total DNA to 4 μg per well) plasmids using Lipofectamine 2000 (Invitrogen) as per the manufacturer's instructions. After 24 h, cells were harvested into reduced serum medium (0.1% FBS) and cultured for an additional 18 h. Wortmannin (200 nM concentration) was added 30 min prior to lysis. Cell lysates were prepared by washing once briefly with PBS, extracting into lysis buffer (20 mM Tris at pH 7.5, 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, 1% Triton X100, 2.5 mM sodium pyrophosphate, 1 mM β-glycerophosphate, 1 mM Na 3 V04, 1 mM DTT, 1 mM PMSF, 10 μg/mL leupeptin, 2 μg/mL pepstatin A, and 10 μg/mL aprotinin) for 10 min at 4 °C, and centrifuging (10 min at 13 500g). Samples were separated by SDS-PAGE and analyzed by immunoblotting with α-BAD pSer112 antibody, mouse α-GST antibody (Cell Signaling Technology #2624), and mouse α-GFP antibody (Clontech #632381) followed by the appropriate secondary antibodies and analyzed on a LI-COR Odyssey CLX infrared imaging system. Signal intensities were quantified using ImageStudio software.

**Cell Death Assays.** COS7 cell lines harboring doxycycline-inducible GFP or GFP-Pim1 (WT or LPVT mutant) cDNA were generated by infection with lentiviruses derived from the pINDUCER20 vector and selection with puromycin. For cell death assays, 3 × 10 5 cells per well were seeded in six-well plates, and the following day, cells were transfected with pcDNA3-HA-BAD 106 (6 μg WT or mutant per well) and empty pcDNA3 (to a final amount of 12 μg DNA per well) using Lipofectamine 2000, according to the standard protocol as per the manufacturer’s instructions. 24 h post-transfection, cells were serum starved (0.1% FBS in growth medium) and treated with doxycycline (500 ng/mL) for 24 h. Cells were harvested by trypsinization, and the proportion of dead cells was determined by staining with 0.4% trypan blue followed by manual counting on a hemocytometer.

**ASSOCIATED CONTENT**

Supporting information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acschembio.7b00089.

Supplementary methods and supporting Figure S1 (PDF)

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**Notes**

The authors declare no competing financial interest.

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