Redesign of the PAK1 Autoinhibitory Domain for Enhanced Stability and Affinity in Biosensor Applications

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The inhibitory switch (IS) domain of p21-activated kinase 1 (PAK1) stabilizes full-length PAK1 in an inactive conformation by binding to the PAK1 kinase domain. Competitive binding of small guanosine triphosphatases to the IS domain disrupts the autoinhibitory interactions and exposes the IS domain binding site on the surface of the kinase domain. To build an affinity reagent that selectively binds the activated state of PAK1, we used molecular modeling to reengineer the isolated IS domain so that it was soluble and stable, did not bind to guanosine triphosphatases and bound more tightly to the PAK1 kinase domain. Three design strategies were tested: in the first and second cases, extension and redesign of the N-terminus were used to expand the hydrophobic core of the domain, and in the third case, the termini were redesigned to be adjacent in space so that the domain could be stabilized by insertion into a loop in a host cyan fluorescent protein (CFP). The best-performing design, called CFP-PACker, was based on the third strategy and bound the kinase domain of PAK1 with an affinity of 400 nM. CFP-PACker binds more tightly to a full-length variant of PAK1 that is stabilized in the “open” state (Kd = 3.3 μM) than to full-length PAK1 in the “closed” state (undetectable affinity), and binding can be monitored with fluorescence by placing an environmentally sensitive fluorescence dye on CFP-PACker adjacent to the binding site.

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Introduction

p21-activated kinase 1 (PAK1) is a serine/threonine kinase that regulates a diverse set of cellular activities including cytoskeletal remodeling, growth factor and steroid receptor signaling, transcription and mitosis.1 The activity of PAK1 is tightly regulated, and the altered expression and activity of PAK1 has been implicated in several human cancers.2 Inactive PAK1 forms an autoinhibited homodimer that is stabilized by interactions between its inhibitory switch (IS) domain and the kinase domain. In the autoinhibited state, residues immediately C-terminal to the IS domain, called the KI segment, occupy the cleft between the N lobe and
the C lobe of the kinase domain and block the active site (Supplementary Fig. S1). PAK1 can be activated by a variety of signaling molecules including the small guanosine triphosphatases (GTPases) Cdc42 and Rac1. In the GTP-bound state, Cdc42 and Rac1 bind to the p21-binding domain (PBD; also known as the Cdc42/Rac-interactive binding (CRIB) domain) of PAK1, which disrupts the autoinhibitory interaction between the IS domain and the kinase domain. The PBD (residues 70–113) and the IS domain (residues 81–136) structurally overlap, and therefore, binding of the GTPase to the PBD is mutually exclusive with binding between the IS domain and the kinase domain. Following GTPase binding, autophosphorylation further stabilizes the activated state.

Because PAK1 activity is under tight spatiotemporal control, we are interested in creating a biosensor that can detect the localization and kinetics of PAK1 activation in living cells. Recently, a PAK1 biosensor (called Pakabi) was created by fusing a yellow fluorescent protein and a cyan fluorescent protein (CFP) to the termini of PAK1. Upon activation and disruption of the interaction between the IS and the kinase domains of PAK1, the distance between the PAK1 termini increases and the FRET (fluorescence resonance energy transfer) signal between the fluorescent proteins decreases. Pakabi has been used to visualize PAK1 activation during cellular spreading and during the formation of cellular protrusions. One limitation of this type of sensor is that it does not detect endogenous PAK1, but rather overexpressed protein. Additionally, to provide a larger FRET signal in the inactive state, the first 64 residues of PAK1 were removed when creating Pakabi. As a result, Pakabi is missing binding sites that localize Pakabi to tyrosine kinase receptors via the adaptors Nck and Grb2. Localization of Pakabi to tyrosine kinase receptors has been shown to be important for actin remodeling, cell motility and the extension of membrane lamellae.

An alternative type of biosensor is one that detects the activation state of an endogenous protein. One strategy for accomplishing this goal is to make use of proteins known as affinity reagents that bind only to the protein of interest when it is in an activated state. Nalbant and coworkers used this strategy to create a biosensor for activated Cdc42. An environmentally sensitive dye was covalently attached to a domain from the Cdc42 effector protein WASP that binds only activated Cdc42. The labeled protein shows a strong increase in fluorescence when binding endogenous, activated Cdc42 and has been used to study the spatiotemporal dynamics of Cdc42 activation in living cells. Here, we rationally design a modified fragment of the PAK1 autoinhibitory domain that can be used similarly to create a biosensor for activation of endogenous PAK1. In many cases biosensors cannot be created because no known domains with appropriate binding characteristics exist; achieving this through the design of novel binding domains or improvement of naturally occurring binders would be important in extending the applicability of biosensors.

Upon PAK1 activation, the interaction between the IS domain and the kinase domain of PAK1 is broken, exposing the IS binding surface on the kinase domain. In previous work, we showed that a protein designed to interact with the IS binding site on the PAK1 kinase domain was sensitive to whether PAK1 was in the “open” or “closed” state. However, the designed protein only bound the open state with an affinity of 100 μM, too weak to be used as the basis for a biosensor. Here, we examine if variants of the IS domain can be used as affinity reagents that differentiate active and inactive PAK. The wild-type IS domain is unfolded in isolation and has only modest affinity for PAK1. To improve the stability of the IS domain and its affinity for PAK1, we used the molecular modeling software Rosetta to identify N- and C-terminal extensions that stabilize the domain and allow for favorable insertion in a host protein. The best-performing redesign binds the PAK1 kinase domain with an affinity of 400 nM and has no measurable affinity for full-length PAK1 when it is in the closed state. We also show that binding can be detected by placing an environmentally sensitive dye adjacent to the binding interface.

Results

The wild-type IS domain is not a good affinity reagent

We first examined if the wild-type IS domain could be used as an affinity reagent that is specific for activated PAK1. We expressed and purified residues 83–137 of PAK1 (WT_IS), which in the crystal structure of autoinhibited PAK1 interact with the C lobe of PAK1 kinase domain and form a small folded domain with three helices and a β-strand (Fig. 1). We did not include residues 138–149 of PAK1, which bind in the active site of the kinase. The circular dichroism (CD) spectrum of the purified IS domain showed that the domain does not fold in isolation (Fig. 2). Not surprisingly, the domain was prone to aggregation and proteolysis during expression and purification. In isothermal titration calorimetry (ITC) experiments, the IS domain bound to the kinase domain of PAK1 with an affinity of only 4 μM, while biosensor-target interactions typically have a dissociation constant of less than 1 μM.
Design strategy

We tested three design strategies for stabilizing the IS domain (Fig. 1a). In all the designs, the goal was to keep the kinase-interacting residues intact while redesigning the surrounding residues to improve stability and destroy binding with Cdc42 and Rac1 (Fig. 1b).

The first strategy we tested was inspired by the crystal structure of PAK1 in its autoinhibited conformation [Protein Data Bank (PDB) code 1F3M]. In this structure, PAK1 forms a homodimer in an autoinhibited conformation with a β-strand from one IS domain interacting with the partner IS domain (magenta). The rest of the partner is not displayed for clarity. (a) Three alternative designs were tested, “IS_hairpin,” “IS_helix,” and “PAcKer.” The new structural elements added to each variant are shown in green. The cyan regions were left largely intact except for some point mutations that allowed better packing with the new structural elements. (b) Multiple sequence alignment of the variants shows that the new structural elements were at the N-terminus or C-terminus, while the residues involved in binding to the kinase domain were left unmutated (black arrows). The residue numbering is based on the primary sequence of full-length PAK1.

![Fig. 1. Structural and sequence comparison of the designs. All the designs were based on the IS domain (IS, cyan), which interacts with the C lobe of the PAK1 kinase domain (yellow). In the crystal structure, PAK1 forms a homodimer in an autoinhibited conformation with a β-strand from one IS domain interacting with the partner IS domain (magenta). The rest of the partner is not displayed for clarity. (a) Three alternative designs were tested, “IS_hairpin,” “IS_helix,” and “PAcKer.” The new structural elements added to each variant are shown in green. The cyan regions were left largely intact except for some point mutations that allowed better packing with the new structural elements. (b) Multiple sequence alignment of the variants shows that the new structural elements were at the N-terminus or C-terminus, while the residues involved in binding to the kinase domain were left unmutated (black arrows). The residue numbering is based on the primary sequence of full-length PAK1.](image-url)
accomplish this by adding additional residues to the protein, rather than mutating preexisting residues in the protein. This model was called IS_helix.

In a third approach, a 16-residue helical stretch was created at the C-terminus of trunc-WT_IS-2 (residues 87–137). The goal of this approach was to bring the N- and C-termini of the domain close to each other so that it can be inserted into a loop in the CFP. The domain was inserted between residues Glu172 and Asp173 of CFP. A small GS linker was added between the N-terminal of the designed domain and Glu172 of CFP. Similarly, another GS linker was added between the C-terminal of the designed domain and Asp173 of CFP. There are two potential benefits of this approach. First, the host protein (CFP) may provide further stability to the IS domain by holding the N- and C-termini adjacent to each other. Second, biosensors often require a stable nonresponsive fluorophore attached to the responsive portion of the biosensor to use as a baseline signal for ratiometric imaging. As with the IS_helix design, the new helix in this case was packed against the rest of the domain and used to expand the hydrophobic core of the domain.

Residues involved in Cdc42 and Rac interactions were also mutated in this design. This design was called PAcKer or CFP-PAcKer when inserted into CFP.

**Design protocol**

We used a computational protocol within the Rosetta molecular modeling program that iterates between structure and sequence optimization to design the new secondary structural elements described here. In brief, the protocol uses three- and nine-residue fragments drawn from the protein database to build loops or extensions of appropriate size. The fragments were picked on the basis of target secondary-structure propensity. During optimization of the helical extensions (IS_helix and PAcKer) or the designed hairpin (IS_hairpin), the new residues were allowed to adopt alternative conformations, while the backbone of the rest of the domain was held fixed. Residues from the core of the domain that packed against the new helical extensions were allowed to adopt alternative side-chain conformations and identities. Distance-based constraints were used to induce the new helical extensions to pack against the rest of the domain.

**Expression and solubility of the designs**

IS domain, IS_hairpin and PAcKer were expressed with an MBP (maltose-binding protein) tag, while IS_helix was expressed with a GST (glutathione S-transferase) tag in *Escherichia coli*. PAcKer inserted into CFP (called CFP-PAcKer) was expressed with a hexahistidine (6×His) tag. Expression levels were highest for IS_helix. Once the proteins were cleaved from GST or MBP, and concentrated, IS_helix showed no signs of aggregation up to a concentration of 1 mM, while wild-type IS and IS_hairpin had lower solubility (~200 μM), and PAcKer showed an intermediate level of solubility.

**Structural characterization of the designs**

Far-UV CD scans showed that IS, IS_hairpin and PAcKer were primarily unfolded, while IS_helix had strong helical content (minima at 208 and 222 nm; Fig. 2a). CD was also used to monitor the thermal denaturation of IS_helix. The midpoint for unfolding was ~35 °C and the curve showed modest cooperativity, as would be expected for a small protein. To further test the ability of IS_helix to form a stable folded structure, we performed a far-UV CD...
scan in the presence of trimethylamine N-oxide (TMAO), an osmolyte that has been shown to stabilize proteins. We saw a considerable increase in helical content at osmolyte concentrations of 0.5 and 1.0 M (Supplementary Fig. S2a), and in the presence of 2 M TMAO, the midpoint for unfolding was 55 °C (Supplementary Fig. S2b).

**Binding affinity and suitability as an affinity reagent**

The binding affinities of IS and the engineered proteins for the PAK1 kinase domain (a model for the active conformation of PAK1), PAK1 full length (a model for autoinhibited PAK1) and PAK1 full length with the mutations V127E and S144E (a model for full-length PAK1 in the open state) were measured with ITC. Val127 stabilizes the interaction between the IS domain and the kinase domain, and mutation to glutamate destabilizes this autoinhibitory interaction in full-length PAK1. We speculated that the mutation S144E will mimic the phosphorylation of Ser144 that has been known to significantly contribute to PAK1 activation. IS_helix, IS_hairpin and wild-type IS (WT_IS) bound to the PAK1 kinase domain with affinities of 1.6, 2.6 and 4 μM, respectively. (Due to low ITC signal-to-noise ratio observed for kinase domain titrated into IS_helix, binding was reconfirmed by fluorescence polarization technique and was found to be 15 μM). It was surprising that IS_helix and WT_IS domain bound with such similar affinities, as the IS_helix was observed to be prefolded, while the WT_IS folds upon binding the kinase. One interpretation of these results is that IS_helix is not preordered correctly, and the structure must rearrange—at some cost to binding energy—to interact with the kinase domain. In this scenario, the reordering energy for IS_helix may be equivalent to the loss in conformational entropy that the IS domain undergoes upon binding.

CFP-PAcKer bound to the PAK1 kinase domain with an affinity of 400 nM, 10-fold tighter than the IS domain (Fig. 3a). It did not bind to closed full-length PAK1 (Fig. 3b), and the binding affinity for the open variant, PAK1(V127E/S144E), was 3.3 μM (Supplementary Fig. S3). In order to test if CFP-PAcKer was binding to the kinase domain using the intended interaction surface, the valine on PAcKer that

![Fig. 3. ITC assays to determine the binding affinities of CFP-PAcKer against active and autoinhibited conformations of PAK1. (a) PAK1 kinase domain (208 μM; model for active conformation) was titrated against 18 μM CFP-PAcKer. An equilibrium dissociation constant of 400 nM was observed with a molar ratio of ~1. (b) PAK1 full length (262 μM; model for “closed” autoinhibited form) was titrated against 21 μM CFP-PAcKer. The titration failed to show any appreciable heat released or absorbed. The data sets were drawn on similar scale for comparison.](image-url)
corresponded to residue 127 in full-length PAK1 and was mutated to a glutamate. As expected, no binding was detected between CFP-PAcKerV127E and the kinase domain. Similarly, mutating Leu470 on the PAK1 kinase domain to glutamate abolished binding between the two proteins (Supplementary Fig. S4). Leu470 is at the center of the interface formed between the IS domain and the kinase domain as seen in the crystal structure (PDB code 1F3M). These results indicate that CFP-PAcKer binds to the target surface area on PAK1 and that binding depends on the conformational state of PAK1.

To create CFP-PAcKer, a helical extension was added to the C-terminus of the IS domain to bring the N- and C-termini in closer proximity. To test if this new structural element was contributing to favorable binding with PAK1, we tested an alternative design in which trunc-WT_IS-2 domain (the starting sequence for PAcKer design) was inserted into CFP, but a glycine–serine linker (GSGSGS to residue 87 and GSGS to residue 137) was used to insert between Glu172 and Asp173 of CFP. This alternative design bound to the PAK1 kinase domain weaker than CFP-PAcKer, 1.65 μM compared to 400 nM (Supplementary Fig. S8).

PAK1 kinase domain induces fluorescence enhancement of a biosensor based on CFP-PAcKer (mero-CFP-Packer)

A lysine in CFP-PAcKer that corresponds to Lys134 in the PAK1 sequence was mutated to a cysteine and covalently modified with an environment-sensitive merocyanine dye designed to respond to protein interaction. This position is adjacent to the predicted interface between CFP-PAcKer and PAK1. We therefore predicted that PAK binding would create a change in the local environment of the dye and lead to a change in fluorescence. We monitored both dye fluorescence (excitation wavelength 593 nm and emission wavelength 620 nm) and CFP fluorescence (excitation at 433 nm and emission at 474 nm) while titrating the biosensor with the kinase domain. The normalized ratio $\frac{[\text{Dye/CFP}]_{\text{Titrant}}}{[\text{Dye/CFP}]_{\text{Buffer}}}$ fit to a "one-site binding" curve gave a binding affinity of 600 nM, consistent with the equilibrium dissociation constant observed in ITC (Fig. 4). There was also a change in normalized fluorescence ratio when titrating with PAK1 V127E/S144E (full-length PAK1 stabilized in the open state). There was no change in normalized fluorescence ratio when mero-CFP-PAcKer was titrated with full-length PAK1 in the autoinhibited state or PAK1 kinaseL470E mutant. Additionally, when dye was conjugated to CFP-PAcKerV127E, there was no change in ratiometric fluorescence when titrating with the PAK1 kinase (Supplementary Fig. S5).

Specificity of the CFP-PAcKer-derived biosensor

Mero-CFP-PAcKer is expected to interact with all group 1 PAKs (PAK 1–3) because the target binding surface on the kinase domain is identical
in all three cases. Group 2 PAKs (PAK 4–6) are more divergent and therefore may bind weaker to CFP-PAcKer. We tested the PAK5 kinase domain, expressed as a GST fusion, for binding to CFP-PAcKer. ITC did not show any appreciable heat absorbed or released when the proteins were mixed (Supplementary Fig. S6) and CFP-PAcKer conjugated to merocyanine fluorescent dye showed only small changes in ratiometric fluorescence in the presence of 1 μM of GST–PAK5 kinase domain (Fig. 5). However, at higher concentrations of GST–PAK5 (>8 μM) the change in normalized fluorescence was larger than with PAK1 (Supplementary Fig. S7). These results indicate that mero-CFP-PAcKer binds weaker to PAK5 than PAK1, but the change in fluorescence is greater when binding to PAK5. In addition to sequence divergence in the binding site, several of the residues surrounding the binding site vary in PAK1 and PAK5 and are expected to create different local environments for the merocyanine dye.

Discussion

We redesigned the IS domain of PAK1 for improved function as an affinity reagent. The focus of the design process was to stabilize tertiary structure formed by the IS domain when it binds the kinase domain of PAK1. In isolation, the IS domain is natively unfolded. The potential advantages of this approach are several: a well-folded domain should be more resistant to proteases, solubility should be improved by burying hydrophobic amino acids in the core of the domain, and preordering the polypeptide in a conformation that is competent for binding should reduce losses in conformational entropy upon binding and increase binding affinity. The designed protein IS_hairpin did not fold in isolation and did not show a strong improvement in solubility. In contrast, IS_helix formed helical structure and was considerably more soluble. However, the redesign did not have enhanced affinity for PAK1, presumably because the structure was not preordered in a conformation appropriate for binding.

Our most successful design, CFP-PAcKer, included a C-terminal extension modeled to bring the N- and C-termini of the IS domain adjacent to each other, which allowed for insertion into a loop in CFP. Inserting the domain into CFP led to increased affinity for PAK1. The rigid structure of the fluorescent protein likely holds the N- and C-termini of the IS domain adjacent to each other and stabilizes the domain in a binding-competent conformation. CFP has high solubility and likely contributed to the enhanced solubility of the fused protein. CFP fluorescence can be used in ratiometric imaging to normalize fluorescence signals generated by merocyanine dyes attached to the IS domain. Instead of designing a helical extension to bring the N- and C-termini of the IS domain close in space, we also examined the use of linkers made from glycine and serine. This design did not show enhanced affinity for PAK1, perhaps because the flexible linkers prevented the preordering of the domain in a binding-competent conformation.

The mero-CFP-PAcKer biosensor has the potential to be a powerful tool for biological studies, shedding light on PAK dynamics in living cells. With a practical affinity reagent in hand, next steps will include examination of different dyes and different dye attachment positions on the biosensor. Preliminary studies in cells showed localized fluorescence increases potentially attributable to PAK activity (data not shown), leading us to move forward with full validation demonstrating selectivity and robust response in vivo. This study demonstrates that a naturally occurring affinity reagent not suitable for biosensor studies can be optimized through rational engineering and modeling, expanding the reach and practicality of live-cell biosensor designs.

Materials and Methods

Modeling using Rosetta

To design IS_hairpin, IS_helix and PAcKer, we used Rosetta’s backbone sampling and design protocols. Using the Robetta server, we created a customized library of fragments from the PDB for each three- and nine-residue window in a fasta sequence. A pseudo-sequence for the region to be modeled was used to bias the fragments to a desired secondary structure. For IS_hairpin, the query fasta sequence consisted of residues 79–84 (SDFEHT) from IS domain chain B (PDB code 1F3M), INGTTI for the canonical hairpin, and residues 86–137 from IS domain chain A (Supplementary Fig. S1a). For IS_helix, the query fasta sequence was a polyalanine sequence, A_{16} with (GS)_{2} sequence for linker followed by residues 100–137 of the IS domain (chain A of PDB 1F3M). The query fasta sequence for PAcKer was created using sequence 87–137 of the IS domain (chain A of PDB 1F3M) followed by GA_{13}SG sequence. The fragments corresponding to the region to be modeled were then assembled using Monte Carlo simulated annealing technique. Fragment insertions were accompanied by a “wobble” operation where the backbone torsion angles (ψ, θ) were perturbed. Monte Carlo minimization protocol was used to iterate between sequence optimization and structure prediction to optimize all-atom energy function in Rosetta. In brief, the function is a linear weighted sum of 12-6 Lennard–Jones potential, the Lazairidis–Karplus implicit solvation model, an orientation-dependent hydrogen-bonding potential, backbone-dependent rotamer probabilities, a knowledge-based electrostatic energy term, and amino acid
The cells were grown to an OD600 of 0.6 with an N-terminus 6×His tag. Protein was expressed as GST fusion with a thrombin cleavage site and cloned in pQE-80L vector so that the MBP fusion with a TEV (tobacco etch virus) protease sequence, IS_hairpin and PAcKer were expressed as an protein.

LYFQSM) in the N-terminus and expressed as a GST fusion with an extra sequence (MHHHHHHSSGVDLGTEN-

extinction coefficient of 44,000 M−1 cm−1 was estimated using CFP absorbance at 433 nm with an and absorbance at 280 nm. CFP-PAcKer concentration was estimated using theoretical molar extinction coefficients (Amicon Ultra, Millipore). Protein concentration was measured with a VP-ITC isothermal titration calorimeter (MicroCal, GE Healthcare). The designs were placed in the cell at concentrations between 14 and 21 μM (IS_helix at a concentration of 50 μM). PAK1 variants were used as titrants at 12- to 15-fold higher concentrations. The proteins were exhaustively dialyzed in 20 mM sodium phosphate buffer (pH 7.4), 25 mM NaCl and 5 mM 2-mercaptoethanol (BME). For each experiment, 29 titrations of 10 μl were made. The data were analyzed with Origin50 software and fit with a model for one-site binding.

Fluorescence polarization assay

Due to a very low signal observed in ITC for IS_helix binding experiment, the binding affinity of IS_helix for PAK1 kinase domain was also measured by fluorescence polarization.28 Ser36 in IS_helix sequence (corresponding to Ser115 in full-length PAK1 sequence and chain A of PDB 1F3M) was mutated to cysteine for conjugation of the thiol-reactive fluorescent probe BODIPY 507/515 iodoacetamide (Molecular Probes). Fluorescence polarization assays were carried out on a Horiba Spex FluoroLog-3 instrument (Jobin Yvon Inc) performed in L-format with the excitation wavelength set at 508 nm and emission wavelength set at 545 nm. BODIPY-conjugated IS_helix [in 50 mM Tris–Cl (pH 7.5), 5 mM BME] at a final concentration of 5 μM and volume of 180 μl was titrated with PAK1 kinase domain [in 50 mM Tris–Cl (pH 7.5), 50 mM NaCl and 5 mM BME].

Conjugation of fluorescent dye to CFP-PAcKer

To enable detection of CFP-PAcKer binding to PAK1, we conjugated a solvent-sensitive merocyanine dye18 to CFP-PAcKer (mero-CFP-PAcKer). Freshly reduced [using 1 mM tris(2-carboxyethyl)phosphine (TCEP)] CFP-PAcKer with L134C (residue numbering based on primary sequence of full-length PAK1 and chain A of PDB 1F3M) mutation was buffer-exchanged to 50 mM sodium phosphate (pH 7.5) with a PD10 desalting column followed by overnight proteolysis using TEV protease and then again loading the product on the Ni-NTA column. WT_IS domain, IS_hairpin and PAcKer were purified using an Ni-NTA purification step (Source 15Q beads, GE Healthcare). All the design variants eluted at similar

Redesign of the PAK1 Autoinhibitory Domain

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column (Amersham Biosciences). Three to five equivalents of merocyanine dye dissolved in DMSO (dimethyl sulfoxide) was added to 1-ml aliquot of protein (typical concentration 40–100 μM) in an Eppendorf tube and immediately mixed by inverting the tube multiple times. The tube was then covered with aluminum foil and left under gentle mixing for 2 h at room temperature. The conjugation reaction was then terminated with an excess of BME (5-μl of 14.3 M stock in 1 ml of reaction mix). Unreacted dye was separated from the conjugated protein with a PD10 column equilibrated and then eluted with 20 mM sodium phosphate (pH 7.4), 25 mM NaCl, and 5 mM BME. Dye-labeled protein concentration was estimated using CFP absorbance at 433 nm (ε = 44,000 M⁻¹ cm⁻¹). Dye concentration was estimated using dye absorbance at 593 nm (ε = 120,000 M⁻¹ cm⁻¹). The conjugation efficiency (dye concentration/CFP concentration) was ~100%.

Fluorimetric measurement of mero-CFP-PAcKer interaction with PAK1

Dye-labeled CFP-PAcKer [200 nM mero-CFP-PAcKer, 20 mM sodium phosphate (pH 7.4), 25 mM NaCl, 5 mM BME] was titrated with PAK1 variants, 0–10 μM. The reaction mix for each titration point was made in a separate Eppendorf tube and incubated for 10 min at room temperature. Readings were taken on a Horiba Spex FluoroLog-3 instrument (Jobin Yvon Inc) with a 3-mm cuvette. The sample was excited at 433 nm (for CFP) and 593 nm (for dye) and the emission signal was measured at 474 nm (CFP signal) and 620 nm (merocyanine dye signal), respectively. The readings were taken from lower titrant concentration to higher without an intermediate cuvette washing step. Data accumulation took 30 min on average from the start (0 titrant reading) to the end of the experiment (10 μM titrant reading). A normalized fluorescence ratio [(Dye/CFP)Titrant / (Dye/CFP)Buffer] was calculated and plotted against titrant concentration. The data was fit using SigmaPlot and Michaelis–Menten equation for one-site binding. Experiments were also conducted with the GST–PAK5 kinase domain in a similar format.

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Supplementary Data

Supplementary data to this article can be found online at doi:10.1016/j.jmb.2011.08.022

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