

# mRNA Display Selection of a High-Affinity, Modification-Specific Phospho-I $\kappa$ B $\alpha$ -Binding Fibronectin

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ABSTRACT The complexity of the human proteome is greatly expanded by post-translational modifications. New tools capable of recognizing these modifications in a sequence-specific fashion provide a route to purify these modified proteins, to alter protein trafficking, and to visualize signal transduction in real time. Here, we have evolved novel, modification-specific ligands that target phosphorylated  $I \kappa B \alpha$ . To do this, we employed mRNA display-based in vitro selection using a 30-trillion-member protein library based on the fibronectin type III domain. The selection yielded one fibronectin molecule, 10C17C25, that binds a phospho-IkB $\alpha$  peptide with  $K_d$  = 18 nM and is over 1000-fold specific compared to the nonphosphorylated peptide. 10C17C25 specifically recognizes endogenous phosphorylated  $I \kappa B \alpha$  from mammalian cell extract and stabilizes phospho-lkB $\alpha$  in vivo. We also incorporated 10C17C25 into a FRET indicator that detects IkB kinase (IKK) activity in vitro, demonstrating the utility of selecting designed adaptors for kinase activity sensors.

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hereas the human genome contains less than 25,000 genes, the proteome likely contains over 1,000,000 different proteins when alternative splicing and post-translational modifications (PTM) are taken into account (1). Posttranslational modifications are essential to cellular signaling, and the ability to detect and quantitate modifications is challenging but immensely valuable. Nucleic-acid-based microarrays are powerful proteomic tools, although they cannot detect the presence and extent of PTMs and may not be accurate predictors of protein quantities (2). Attempts to generate open-access resources for modification-specific affinity reagents to all proteins are currently under way (3). In addition to detection in vitro, novel protein affinity reagents that are functional in the intracellular environment are also useful for determining protein localization and for assessing protein function (4, 5). Compared to gene and transcript knockout techniques, domain and state-specific protein binders that knock down activity may be better determinants of protein function (6). Increasingly, in vitro selection techniques such as ribosome display and mRNA display are being implemented to generate novel proteinbased affinity reagents with alternative scaffolds (7–9). Nonimmunoglobulin scaffolds such as the ankyrin-based "DARPins" (10) and fibronectin type III "monobodies" (11) have advantages over antibodies for both in *vitro* and *in vivo* applications because they (i) lack disulfide bonds, (ii) can be expressed efficiently in bacteria and eukaryotic cells, and (iii) possess improved stability (12).

Intrinsically unstructured domains compose a large fraction of the proteome, and many are subjected to PTM (1, 13). An example of such a domain is the N-terminus of IκB. In the classical NF-κB pathway, three IκB proteins, IκBα, β, and ε, regulate the activity of the ubiquitous transcription factor NF-κB (14). NF-κB plays an important role in inflammation and is implicated in autoimmune disease and cancer (15). Upon stimulation, IkB kinase (IKK) immediately phosphorylates  $l \kappa B \alpha$  at serines 32 and 36 (16, 17).  $I\kappa B\alpha$  is then ubiquitinated by the SCFβTrCP E3 ligase complex and degraded by the proteasome, thereby freeing NF-κB to direct transcription (18). Reagents that detect or inhibit the degradation of  $l\kappa B\alpha$  in vivo may be useful in order to understand the many complex pathways to NF-κB activation. Inhibition at this control point is ideal since the functions of IKK and βTrCP are not specific to the classical NF-κB pathway (15, 19).

We employed *in vitro* selection by mRNA display to generate novel protein affinity reagents that recognize phosphorylated  $I\kappa B\alpha$  specifically. We utilized a combinatorial protein library based on the 10th fibronectin type III domain of human fibronectin (10FnIII) to generate protein molecules that are functional both *in vitro* and *in vivo* (20). The target used for this selection was a synthetic phospho-peptide corresponding to

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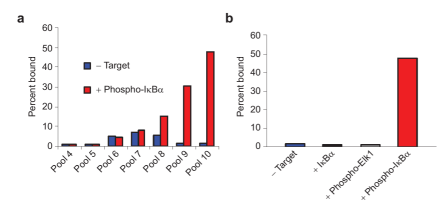


Figure 1. Selection of phospho- $l\kappa B\alpha$  binders. a) Enrichment of target binders was monitored by radiolabeled fusion binding assays. b) Pool 10 binding is sequence-specific and phosphospecific.

amino acid residues 22-41 of human  $I\kappa B\alpha$ . This peptide contains the E3 ligase recognition sequence including phosphorylated serines 32 and 36, which are essential for degradation. The target peptide contained an N-terminal biotin spaced by aminohexanoic acid for specific immobilization to either neutravidin agarose or streptavidin acrylamide beads.

The first round of selection was carried out so that the first affinity enrichment step contained at least three copies of the 30trillion-member 10FnIII library (21). Enrichment was monitored by measuring radiolabeled fusion binding (Figure 1, panel a). In order to reduce the enrichment of bead binders, a negative selection step was employed at round 6. After 10 rounds of selection, pool 10 was determined to have approximately 50% binding efficiency at RT with relatively low background binding (1%). Figure 1 (panel b) demonstrates that pool 10 binding is both sequence-specific and phospho-specific. Binding to a nonphosphorylated  $l\kappa B\alpha$  peptide otherwise identical to the target was equal to background. Likewise, phospho-serines alone are not sufficient for binding. No binding was detected above background to a peptide corresponding to Elk1 residues

380—394 containing two phosphorylated serines.

Pool 10 was cloned, and 11 sequences were obtained. All 11 sequences were highly similar, suggesting that all sequences share a common ancestor (see Supporting Information). One representative sequence, 10C17, is illustrated in Figure 2, panel a. The wild-type sequence shown represents a 7-residue deletion mutant previously described as 10FnIII ( $\Delta 1-7$ ) and has equivalent expression, solubility, and folding stability as full-length 10FnIII (20). Essentially, only one solution to the molecular recognition of phospho-l $\kappa$ B $\alpha$  was obtained. This may be due to an intrinsic rarity of binders in the library, as obtaining binders to nonstructured targets may be more difficult than obtaining binders to structured targets. Alternatively, it is possible that the pool reached an artificial bottleneck due to overly stringent selection conditions or that the immobilization scheme on streptavidin beads limited accessibility of the target.

The ability for the binders to express in bacteria is important for their utility as affinity reagents. Both 10C17 and another pool 10 clone, 10C19, do not express well in *E. coli* BL21(DE3) (data not shown). In order to improve the expression of 10C17, solubility

was evolved using the GFP reporter screen developed by Waldo et al. (22). Three rounds of evolution (A, B, and C) were performed using error-prone PCR so that one or two mutations were generated per gene. Representative sequences obtained from each of the three rounds of evolution performed are listed in Figure 2, panel a. Approximately 2000 colonies were screened in each round, and the brightest 24 colonies were picked and assayed. The improvement in expression was analyzed by comparing total cell fluorescence of 10C17-GFP variants relative to that of WT 10FnIII (Figure 2, panel b). For the first two rounds of evolution, expression was induced with IPTG for 4 h (screen 1). We found that although the initial mutations improved expression, the result of the long induction was that fluorescent aggregates were able to accumulate. This resulted in obtaining variants that performed better than wildtype (WT) 10FnIII in the assay but did not express as well. We found that decreasing the length of expression to 1.5 h improved the sensitivity of the screen (Figure 2, panel b, screen 2). In order to ensure that function was retained, binding was measured for each of the variants using SDS-PAGE quantitated by autoradiography (Figure 2, panel c).

From round one of the evolution, one variant containing two mutations was obtained (A21) that improved both solubility and binding (Figure 2). Two sequences were obtained from round two that improved solubility and did not affect binding. B11 was a unique clone, whereas the B01 sequence was found three times out of the 24 clones assayed. We created 10C17B25 by combining the two beneficial point mutations selected, L13Q and A10T, from B11 and B01, respectively. 10C17B25 displays an additive improvement in solubility. The third round of evolution again generated two variants both with improved expression (C21, which was found twice, and C22, which was

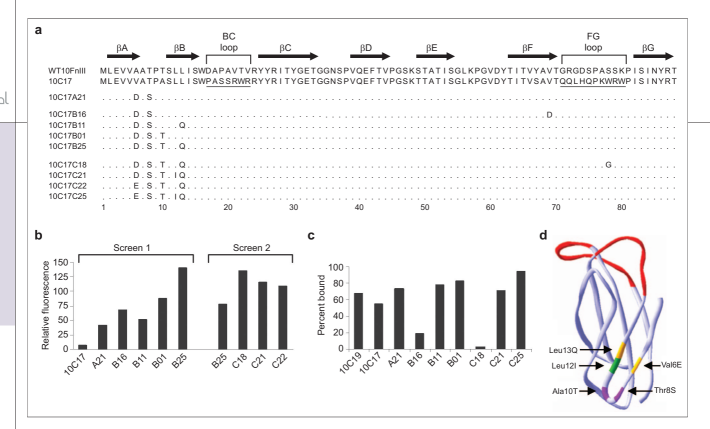


Figure 2. Solubility evolution of selection winner using a GFP reporter screen (22). a) Sequence of wild-type 10FnIII, a pool 10 clone (10C17), and evolved variants of 10C17 from three rounds of evolution. Dots represent sequence identical to 10C17. b) Total cell fluorescence of expression cultures relative to wild-type 10FnIII. Proteins were expressed for 4 h (screen 1) or 1.5 h (screen 2). c) Binding of selected clones with *in vitro* translated protein measured by SDS-PAGE autoradiography (representative data). d) Locations of evolved mutations on the 10FnIII surface include solvent-exposed (yellow), buried (green), and AB loop (purple).

unique). The two mutations selected in C21 and C22 were combined to produce 10C17C25, which binds with approximately 94% efficiency (Figure 2, panels a and c). This variant could be expressed at modest levels in bacteria (1 mg  $L^{-1}$  culture) and was purified by nickel affinity and ion exchange chromatography.

The fact that we obtained improvements in both solubility and binding argues that the mutations increased the overall stability of the domain. Of the six mutations we implemented to obtain the 10C17C25 variant, only four positions differ from WT 10FnIII. Two mutations were obtained at position Val6 (originally Ala in WT 10FnIII), V6D and D6E. Also, Ala10 reverted back to Thr, the wild-type residue (Figure 2, panel a). Mutations V6E and L13Q are notable because they are both likely solvent-exposed and close in space (Figure 2, panel d) (23). At these two positions, the mutations altered what had been a large hydrophobic patch on the protein surface, making it more polar with residues that have the potential to interact favorably. In addition to these

positive mutations, we also found changes that improved solubility but negatively affected function (for example, variants B16 and C18, which were found three and six times out of 24 clones in pools B and C, respectively).

We next addressed the affinity and specificity of 10C17C25, as well as its ability to bind endogenous phospho-lκBα from human cell culture (Figure 3). 10C17C25 was subcloned into a modified pET 21 vector that generates a C-terminal MBP fusion with a C-terminal His-tag. MBP was used to increase the expression of 10C17C25 (45 mg  $L^{-1}$  culture) and provide an additional affinity purification tag. Binding to both the phosphorylated and nonphosphorylated  $I\kappa B\alpha$  peptides was measured using a Biacore T100 instrument (Figure 3, panels a and b). 10C17C25-MBP binds phospho- $I\kappa B\alpha$  with  $K_d = 18$  nM, which is comparable to high-affinity antibodies (24). Binding to the nonphosphorylated peptide was not detected at the highest concentration of 10C17C25-MBP tested (28 µM) (Figure 3, panel b). Therefore, the specificity of

10C17C25 for the phosphorylated state is >1000-fold.

We next wanted to determine whether 10C17C25 is able to recognize wild-type, full-length  $I\kappa B\alpha$  specifically in its phosphorylated state. Phospho- $l\kappa B\alpha$  was produced in 293T cells by stimulation with TNF- $\alpha$  and was stabilized by addition of the protease inhibitor calyculin A and the proteasome inhibitor MG132. Both stimulated and unstimulated cells were lysed with Tris-buffered saline plus 1% triton x-100. 10C17C25-MBP was added to the cleared lysates and incubated with nickel-NTA beads. A Western blot performed to detect the presence of  $I\kappa B\alpha$ using a polyclonal antibody demonstrates that 10C17C25 is able to bind full-length  $I\kappa B\alpha$  only from TNF- $\alpha$  stimulated cells (Figure 3, panel c).  $I\kappa B\alpha$  that is immobilized by 10C17C25 has a noticeable reduction in mobility typical of phosphorylated  $I\kappa B\alpha$ 

We also tested the ability of 10C17C25 to express and function in human cells. 10C17C25 was cloned into a modified pIRES-puro vector containing a C-terminal

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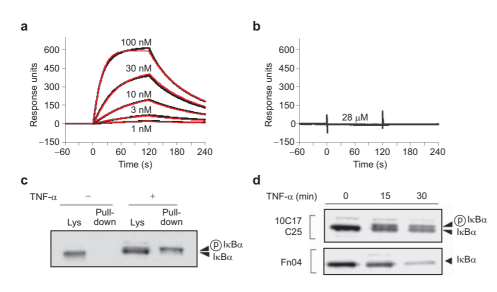


Figure 3. Affinity and specificity of 10C17C25. SPR was performed to test affinity and specificity to phospho-I $\kappa$ B $\alpha$ . Concentrations of 10C17C25-MBP ranged from 1–100 nM for phospho-I $\kappa$ B $\alpha$  binding (panel a) and up to 28  $\mu$ M for nonphosphorylated I $\kappa$ B $\alpha$  peptide binding (panel b). Black represents raw data, red represents curve fits. c) Modification-specific binding to full-length I $\kappa$ B $\alpha$  was determined by pull-down from mammalian cell culture. d) Intracellular function was demonstrated by the inhibition of I $\kappa$ B $\alpha$  degradation in TNF- $\alpha$  stimulated 293T cells transiently transfected with 10C17C25-GFP. Cells transiently transfected with nonselected Fn04-GFP were used as a control.

GFP. This fusion protein was well-expressed throughout the cytoplasm and nucleus of 293T cells when transiently transfected. 10C17C25-GFP is able to stabilize phospholkB $\alpha$  in vivo when transiently expressed (Figure 3, panel d). Expression of 10C17C25-GFP stabilizes phospholkB $\alpha$  even after 30 min of TNF- $\alpha$  incubation, when lkB $\alpha$  degradation is maximal in 293T cells. PhospholkB $\alpha$  is not detected in cells expressing Fn04, a nonselected fibronectin control, as phospholkB $\alpha$  is rapidly degraded without an inhibitor present (Figure 3, panel d).

FRET sensors have been employed to detect PTMs reversibly *in vivo* in a spatial and temporal manner (25–27). The sensors previously developed utilize natural modification-specific binding domains such as SH2 and a chromodomain. These sensors contain an enzyme substrate sequence that is bound by the proximal binding domain upon modification when signaling is active.

This binding restrains the sensor, resulting in a change in FRET between an N-terminal CFP and a C-terminal YFP. The potential for such sensors can be greatly expanded by applying in vitro selection to create novel modification-specific binding domains. We demonstrated this by creating a FRET-based sensor for IKK activity and assessed its function in vitro (Figure 4, panel a). CFP, YFP, and a new multiple cloning site that contains a flexible linker sequence were cloned into a pET21 (His)<sub>6</sub>-tag vector (see Supporting Information). 10C17C25 and a Klenow extended fragment encoding  $l\kappa B\alpha$  residues 23-41 were cloned into the vector in two orientations with short and long linkers (Figure 4, panel b). The second orientation (IKK FS C25 2S and 2L) contains longer linkers than the first (IKK FS C25 1S and 1L) due to the location of the binding surface relative to the termini of the fibronectin domain. A nonselected fibronectin variant was also cloned

into one orientation as a control (IKK FS Fn04 control).

The five constructs were expressed in E. coli BL21(DE3) and purified by nickel-NTA affinity chromatography. The constructs were phosphorylated in vitro with recombinant IKK-β, and the reactions were terminated by addition of EDTA after 1 h. FRET was then measured on constructs with or without incubation with IKK-B (Figure 4, panel c). FRET was quantitated by measuring the ratio of the YFP emission intensity and the CFP emission intensity after excitation at  $\lambda = 435$  nm and is represented as a percent change after incubation with kinase (Figure 4, panel b). The first orientation results in a small decrease in FRET after IKK-β incubation, whereas the second orientation results in a substantial increase in FRET. The length of

the linker does not significantly affect the change in FRET, and phosphorylation of the control sensor does not result in a change in FRET. This indicates that neither phosphorylation alone nor interaction with IKK-β can account for the changes in FRET observed with IKK FS C25 2S and 2L. Likewise, the FRET response is demonstrated to be a result of binding to the phosphorylated  $l\kappa B\alpha$ sequence as excess phospho-peptide reduces the FRET response achieved after incubation with kinase (Figure 4, panel c). Western blot was performed using a monoclonal phospho(S32/S36)-specific anti- $I\kappa B\alpha$  antibody to demonstrate that the kinase reaction is efficient and phosphorylation does occur (Figure 4, panel d).

In conclusion, we have demonstrated the utility of mRNA display using a combinatorial protein library based on the 10FnIII scaffold for modification-specific recognition of protein domains. We have demonstrated that the selection, combined with



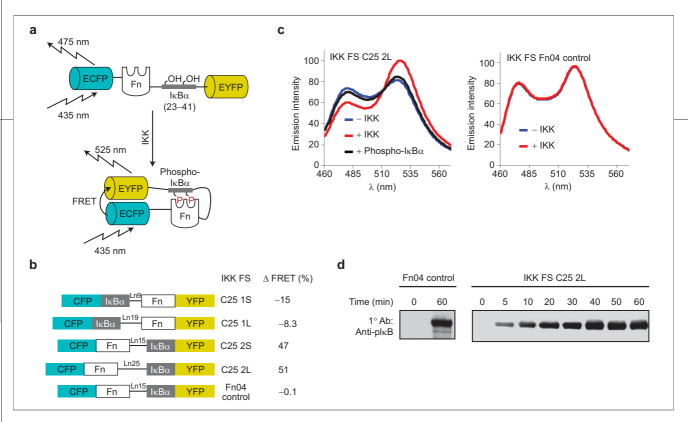


Figure 4. IKK FRET sensor. a) Illustration of a FRET sensor for IKK activity using a designed modification-specific fibronectin (Fn) binder. b) Four sensors were created, as well as a control containing nonselected Fn04. The IKK substrate sequence corresponds to  $I\kappa B\alpha$  residues 23–41. c) Representative fluorescence emission spectra of IKK FRET sensors. d) Western blot to demonstrate both the control and FRET sensor 2L are efficiently phosphorylated by IKK.

solubility evolution, was able to produce a high-affinity, high-specificity binder to phospho- $I\kappa B\alpha$ . This scaffolded library is ideally suited to discrimination of modifications due to the flexible, continuous ligand binding surface generated by the randomized loops (20). We have also demonstrated utility of the library for binding unstructured epitopes with high affinity. 10C17C25 has been demonstrated to not only recognize phospho-lkB $\alpha$  in vitro but also to bind and inhibit degradation in vivo. This intracellular inhibition may prove useful for studying the complex NF-кВ pathway as an alternative to gene knockout or overexpression of modified IkB.

#### **METHODS**

Reagents. Oligonucleotides were synthesized by IDT or the Yale Keck oligonucleotide synthesis facility. Oligonucleotide sequences are listed in Supporting Information. Peptides were synthesized by Biomer-Tech and purified by reverse-phase HPLC. Products were confirmed by electrospray mass spectrometry. For plasmid construction and cloning, see Supporting Information. *E. coli* BL21(DE3) was purchased from Novagen. 293T cells were cultured in Dulbecco's modified Eagle's medium plus 10% fetal bovine serum (Invitrogen).

**Selection.** The library used in this selection was previously described (20). Protein-mRNA fusions were generated as previously described (21).

Briefly, each round of selection includes PCR. in vitro transcription, splint mediated ligation to a 3' puromycin-containing DNA linker (T4 DNA ligase, NEB), urea PAGE purification, reticulocyte lysate translation (Red Nova, Novagen), salt-induced fusion formation, oligo-dT cellulose purification (NEB), reverse transcription (Superscript II, Invitrogen), and affinity enrichment. For the first round of selection, the 30-trillion-member library was generated with at least three copies of fusions. Affinity enrichment was typically performed with 10-20 μL of beads saturated with peptide (neutravidin agarose or streptavidin ultralink beads, Pierce). The selection buffer included PBS supplemented with 0.5 mg mL<sup>-1</sup> BSA (Sigma), 0.1 mg  ${\rm mL^{-1}}$  sheared salmon sperm DNA (Sigma), 0.02% Tween 20, and 1 mM DTT. A negative selection step was performed beginning at round 6 following reverse transcription by flowing fusions over  $\sim$ 100  $\mu$ L of empty blocked beads in column format. Free neutravidin or streptavidin (Pierce) was also including in the selection binding buffer to compete with background binders. Binding was measured by scintillation counting of 35Smethionine labeled, oligo-dT cellulose purified fusions. All radiolabeled binding assays were performed at RT. Pool 10 was cloned into a TOPO-TA vector (Invitrogen) for sequencing.

**Solubility Evolution.** Error-prone PCR was performed with 7 mM MgCl $_2$  and 0.05 mM MnCl $_2$  to generate one or two mutations per gene. The GFP reporter screen and fluorescence quantitation was carried out as previously described (20, 22). In each round, 24 colonies were picked and assayed prior to sequencing. In some cases, three or more identical clones had been picked, and these independent experiments led to standard deviations of 4.1%, 4.0%, and 3.3% for B01, B16, and C18, respectively. To test retention of function, crude

<sup>35</sup>S-Methionine labeled, *in vitro* translated protein was bound to immobilized target peptide or empty beads, washed, and eluted with SDS PAGE loading buffer. An equivalent amount of crude translation product, and eluted protein was analyzed by SDS-PAGE autoradiography to determine percent protein bound.

Surface Plasmon Resonance. SPR was performed on a Biacore T100 instrument (USC NanoBiophysics Core Facility). 10C17C25-MBP was expressed in *E. coli* BL21(DE3) by induction with 1 mM IPTG at mid-log phase for 3 h. Pelleted cells were lysed with B-PER (Pierce) and affinity-purified with a nickel-NTA column (Qiagen), followed by affinity purification with an amylose resin column (NEB) and desalting (PD-10, Amersham Biosciences). Biacore buffer included PBS plus 0.5 mg mL $^{-1}$  BSA and 0.05% Tween 20. Various concentrations of analyte were flowed at  $100~\mu$ L/min for 2 min at RT and allowed to dissociate for 45 min. Binding sensograms were fitted with Biacore evaluation software.

Pull-Down. Confluent 293T cell culture was incubated with TNF- $\alpha$  (20 ng mL<sup>-1</sup>), MG132 (10  $\mu$ M), and calyculin A (50 nM), respectively, for 15 min (Calbiochem). Stimulated and unstimulated cells were lysed for 5 min with ice-cold lysis buffer (PBS, 1% triton x-100, 20 ng mL<sup>-1</sup> MG132, 100 nM calyculin A, protease inhibitor cocktail III (Calbiochem), and 1 mM EDTA). MG132 was required during lysis and binding to prevent degradation and depletion of phosphorylated  $I\kappa B\alpha$ . Cell extracts were cleared by centrifugation at 20,000q for 5 min, combined with purified 10C17C25-MBP, and incubated with nickel-NTA beads for 1 h at 4 °C. After three washes, samples were eluted with PAGE loading buffer. Crude cell extracts from stimulated and unstimulated cells were run with the eluted samples on SDS-PAGE (NuPAGE system, Invitro-

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gen), transferred to PVDF membrane (Biorad), and incubated with polyclonal anti-IkB $\alpha$  (C21) anti-body (Santa Cruz Biotechnology). After secondary antibody incubation, blots were incubated with SuperSignal West Dura substrate (Pierce) and imaged by chemiluminescence with an Alpha Imager equipped with a cooled CCD camera.

In Vivo Inhibition. 293T cells were transiently transfected with 10C17C25-GFP or Fn04-GFP using Lipofectamine Plus reagent (Invitrogen) following the manufacturers recommendations. One day post-transfection, cells were stimulated with 20 ng mL $^{-1}$  TNF- $\alpha$  for the indicated time. Crude cell extracts were prepared and analyzed by Western blot.

IKK FRET Sensor. FRET sensor proteins were expressed as described for 10C17C25-MBP. Nickel-NTA affinity purified sensors (50  $\mu g$  mL $^{-1}$ ) were kinased with recombinant IKKβ (1  $\mu g$  mL $^{-1}$ , Invitrogen) in 20 mM Tris, pH 7.5 plus 10 mM MgCl $_2$  and 200  $\mu M$  ATP. Reactions were terminated by addition of EDTA. Fluorescence emission spectra were obtained using a Shimadzu fluorimeter RF-5301PC by excitation at 435 nm with 5 nm slit widths. FRET was calculated as the emission intensity at 525 nm divided by the intensity at 475 nm. Western blot was performed as described using a monoclonal phospho-lkB $\alpha$  (5A5) antibody (Cell signaling Technology).

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Supporting Information Available: This material is free of charge via the Internet.

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### **Supporting Information**

All oligonucleotides were purchased from Integrated DNA Technologies unless otherwise stated. All oligos longer than 35 nucleotides were purified by urea PAGE. Creation of the 30 trillion member Fn library was described previously (1). Fnoligo1 and Fnoligo8 were used for amplification during selection.

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Fnoligo1: 5'-TTCTAATACGACTCACTATAGGGACAATTACTATTTACAATTA-
CAATGCTCGAGGTCGTCG
Fnoligo8: 5'-GGAGCCGCTACCGGATCCGGTGCGGTAGTTGATGGAGATCGG
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Fnoligo1 contains the T7 RNA polymerase promoter sequence, a portion of the TMV translation enhancer sequence, and anneals to the 5' end of the Fn library. Fnoligo8 anneals to the 3' end of the Fn library, contains a BamHI restriction site, and codes for a (Gly-Ser)<sub>3</sub> spacer. After *in vitro* transcription, mRNA pools were ligated to pF30P (Yale Keck Oligonucleotide synthesis facility) using Fn-pF30P-Splint.

```
pF30P: 5'-phospho-A_{21}-9_3-ACC-Pu 9 = \text{phosphoramidite spacer } 9, \text{ Pu = Puromycin (Glen Research)} \text{Fn-pF30P-Splint: } 5'\text{-TTTTTTTTTTTTGGAGCCGCTACC}
```

The pool 10 PCR product was cloned into pCR4-TOPO TA cloning vector (Invitrogen). Eleven sequences were obtained:

#### Pool 10 sequences:

WT10FnIII MLEVVAATPTSLLISWDAPAVT	TVRYYRITYGETGGNSPVQEFTVPGSKSTATISGLK	PGVDYTITVYAVTGRGDSPASSKPISINYRT
FnIkB10C19PASSRV	WR	QQLRQPKWRW
FnIkB10C08PASSRV	NR	L.QQLRQPKWRW
FnIkB10C04PTSSRV	WR	QQLRQPKWRW
FnIkB10C03PASSRV	NR	CQQLHQPKWRW
FnIkB10C05PASSRV	WRGKK	CQQLHQPKWRW
FnIkB10C16	WR	CQQLHQPKWRW
FnIkB10C18	WR	CCQQLHQPKWRW
FnIkB10C20	WRCDR	CQQLHQPKWRW
FnIkB10C01QPASSRV	WRH	CQQLHQPKWRW
FnIkB10C09VPASSRV	WR	CQQLHQPKWRW
FnIkB10C17VAPASSRV	WRT	SQQLHQPKWRW

Fnoligo1 and Fnoligo8 were used to amplify 10C17 and 10C19 for *in vitro* transcription followed by *in vitro* translation to determine binding efficiencies.

All restriction enzymes and T4 DNA ligase were purchased from NEB. 10C17 and 10C19 were amplified with primers "Fnlibrary5Nhe" and "Fnlibrary3His" for cloning into pET11a.

Fnlibrary5NdeI: 5'-CTATTTACAATTCATATGCTCGAGGTCGTCG

Fnlibrary3His: 5′-GGTTGGTGATCAGTGGTGATGGTGATGGTGATGGGATCCGGTGCGTAGTTG

Fnlibrary5NdeI contains an NdeI digestion sequence and anneals to the 5′ end of the Fn library.

Fnlibrary3His anneals to the 3′ end of the Fn library, contains a BamHI restriction site, codes for a (His)<sub>6</sub>-tag, and contains a BcII restriction site. The PCR products were digested with NdeI and BcII, purified (QIaquick, Qiagen), and were ligated into the NdeI and BamHI restriction sites of pET11a, creating vectors pAO5-10C17 and pAO5-10C19. Insertion of genes into the NdeI and BamHI restriction sites of the new vectors will result in addition of a C-terminal (His)<sub>6</sub>-tag immediately followed by a stop codon. pAO5-10C17 and pAO5-10C19 were expressed in E.

Coli BL21(DE3) but did not produce detectable soluble protein. 10C17-(His)<sub>6</sub> was purified under denaturing conditions but attempts to refold the protein were unsuccessful.

10C17 and subsequent variants were amplified under error prone conditions (7 mM MgCl<sub>2</sub>, 0.05 mM MnCl<sub>2</sub>) using Fnlibrary5NdeI primer and Fnoligo8. The resulting PCR products were digested with NdeI and BamHI and cloned into the GFP-reporter vector generously provided by Dr. Geoffry S. Waldo (2). To test binding, selected variants were amplified with Fnoligo1 and Fnoligo8 for subsequent *in vitro* transcription and translation. 10C17B25 (see Figure 2) was created by amplifying 10C17B11 with 10C17oligoA10T followed by FnoligoNdeI.

10C17oligoA10T: CATATGCTCGAGGTCGTCGATGCATCTCCAACCAGCCTCCAGATCAGC
10C17B25 was used as the template for the third round of solubility evolution. 10C17C25 was created by amplifying 10C17C21 with 10C17oligoD6E.

The gene inserts from selected clones were digested directly from the minipreped GFP reporter plasmids using NdeI and BamHI, purified by agarose gel electrophoresis (QIAquick, Qiagen) and ligated into the NheI and BamHI sites of pAO5 for expression of C-terminal His-tagged proteins lacking GFP.

In order to increase the yields of soluble 10C17C25, two fusion vectors were created with MBP as an N-terminal (pAO7) or C-terminal fusion (pAO9). Both orientations were tested in order to determine if the enhancement in expression mediated by MBP is dependent on fusion orientation. For pAO7, MBP was amplified from pDW363C (3) in two steps using oligos MBP5N1 and MBP3N1 followed by MBP5N2 and MBP3N2.

MBP5N1: 5'-CATCATCACCACGGTACCAGCAAAATCGAAGAAGGTAAACTGGTAATCTGG

MBP3N1: 5'-CTCGAGACGCCGTCGATGGAGCTCGTCGAGCCATTAGTCTGCGCGTCTTTCAGG

MBP5N2: 5'-GAGTAGGCTTCACATATGCATCACCATCATCACCACGGTACCAGC

MBP3N2: 5'-CATTACGTCGACCTAGGATCCACTAGTAAGCTTCTCGAGACGGCCGTCGATGG

The final PCR product was digested with NdeI and SalI. The modified pET28 vector containing the GFP reporter (2) was digested with NdeI and XhoI and agarose gel purified (QIaquick, Qiagen). Ligation of the digested PCR product into the modified pET28 vector results in a vector containing a (His)<sub>6</sub>-tag, MBP, a peptide spacing linker, a factor Xa cleavage site, followed by a multiple cloning site (Figure S1).

# pAO7: N-terminal MBP fusion From pET 28 (Kan) His-Tag → MBP solubility/affinity tag → fXa → MCS → STOP BgIII T7 promoter Xbal lac operator ...AGATCTCGATCCCGCGAAATTAATACGACTCACTATAGGGGAATTGTGAGCGGATAACAATTCCCCTCTAGAAAT Ndel His-Tag Kpnl AATTTTGTTTAACTTTAAGAAGGAGATATACATATGCATCACCATCACCACGGTACCAGCAAAATCGAAGAA MetHisHisHisHisHisGlyThrSerLysIleGluGlu Sacl Xhol HinDIII GGTAAA...1065bp...AAAGACGCGCAGACTAATGGCTCGACGAGCTCCATCGACGGCCGTCTCGAGAAGCTT GlyLys....355aa...LysAspAlaGlnThrAsnGlySerThrSerSerIleAspGlyArgLueGluLysLeu Spel BamHI ACTAGTGGATCCTAGGTCGAGCACCACCACCACCACCACTGAGATCCGGCTGCTAACAAAGCCCGAAAGGAAGCT ThrSerGlySerEnd T7 Terminator GAGTTGGCTGCCACCGCTGAGCAATAACTAGCATAACCCCTTGGGGCCTCTAAACGGGTCTTGAGGGGTTT...

Figure S1. pAO7 vector.

For pAO9, MBP was amplified in two steps with MBP5C1 and MBP3C followed by MBP5C2 and MBP3C.

MBP5C1: ACGACGATAAGGGGAGCTCCAAAATCGAAGAAGGTAAACTGGTAATCTGG

MBP3C: GATTAGCTCGAGGCCATTAGTCTGCGCGTCTTTCAGG

MBP5C2: CTAAGCGGATCCGATTACAAGGATGACGACGATAAGGGGAGCTCC

The final PCR product and the pET28-GFP reporter were digested with BamHI and XhoI for ligation. The vector used for this ligation is identical to the one described by Waldo et al. except that the original multiple cloning site was replaced (See figure S3). The resulting pAO9 vector contains the multiple cloning site followed by a Flag-tag, enterokinase cut site, MBP, and (His)<sub>6</sub>-tag figure S2.

## pAO9: C-terminal MBP fusion From pET28 vector (Kan) MCS → FLAG-Tag/Enterokinase Cleavage site → MBP solubility/affinity tag → His-Tag → STOP BgIII T7 promoter Xbal lac operator ...AGATCTCGATCCCGCGAAATTAATACGACTCACTATAGGGGAATTGTGAGCGGATAACAATTCCCCTCTAGAAAT Ndel Nhel HinDIII BamHI Spel AATTTTGTTTAACTTTAAGAAGGAGATATACATATGGCTAGCACTAGTTGAGTAAGCTTGGATCCGATTACAAG MetAlaSerThrSer-----LysLeuGlySerAspTyrLys Flag-Tag Sacl GATGACGACGATAAGGGGAGCTCCAAAATCGAAGAAGGTAAA...1065bp...AAAGACGCGCAGACTAATGGCCTCGAG AspAspAspAspLysGlySerSerLysIleGluGluGlyLys... 355aa...LysAspAlaGlnThrAsnGlyLeuGlu enterokinase CACCACCACCACCACTGAGATCCGCTGCTAACAAAGCCCGAAAGGAAGCTGAGTTGGCTGCTGCCACCGCTGA HisHisHisHisHisEnd **T7 Terminator** GCAATAACTAGCATAACCCCTTGGGGCCTCTAAACGGGTCTTGAGGGGTTTTTTG...

Figure S2. pAO9 vector.

pAO5-10C17C25 plasmid was digested with either XhoI and BamHI or NdeI and BamHI and the 10C17C25 fragments were agarose gel purified. These fragments were ligated into purified pAO7 or pAO9 digested with XhoI and BamHI or NdeI and BamHI respectively. We found that the orientation of the MBP fusion does not affect the amount of soluble protein expressed and chose pAO9-10C17C25 for large-scale expression.

For mammalian cell culture expression, 10C17C25-GFP was amplified in two steps (from the GFP reporter vector) with Fn5pIRES and GFP3FLAG1 followed by amplification with Fn5pIRES and GFP3FLAG2.

Fn5pIRES: 5'-CTTCTAGCGGCCGCCACCATGCTCGAGGTCGTCG

GFP3FLAG1: 5'-CGTCCTTGTAGTCACCAGAGCCTTTGTAGAGCTCATCCATGCCATGTG

GFP3FLAG2: 5'-GTGACCTGATCACTTATCGTCATCGTCCTTGTAGTCACCAGAGC

Fn5pIRES contains a Not I restriction site, encodes a Kozak sequence, and anneals to the 5' end of the Fn library. GFP3FLAG1 anneals to the 3' end of GPF and encodes a Flag sequence.

GFP3FLAG2 extends the Flag sequence and add a BcII restriction site. The PCR products were digested with NotI and BcII and cloned into the NotI and BamHI sites of pIRESpuro (Clontech).

In order to create a new FRET sensor vector with a versatile multiple cloning site, we obtained ECFP and EYFP from Dr. Jose Aberola-IIa. The ECFP and EYFP clones contain the A207K mutation which eliminates dimerization. ECFP was amplified in two steps with ECFP5NdeI and ECFP3-1 followed by ECFP3-2. EYFP was amplified with EYFP5BamHI and EYFP3SalI.

ECFP5NdeI: 5'-GGAGATATACATATGGTGAGCAAGGGCGAG

ECFP3-1: 5'-CCTCCGGCCGAAGAATTCAAGCTTGAGCTCTTGTACAGCTCGTCCATGCCGAG

ECFP3-2: 5-CATGGAGGATCCGCCGCTCGAGCCGTTGCTGCCTCCGGCCGAAGAATTCAAG

EYFP5BamHI: 5'-AGCGGCGGATCCTCCATGGTGAGCAAGGGCGAGG

EYFP3SalI: 5'-TCGTCGGTCGACCTTGTACAGCTCGTCCATGCCCAGAG

### pKC1: FRET sensor vector From pET 21 (Amp) ECFP(A207K) → MCS/LINKER → EYFP(A207K) → His-Tag → STOP T7 promoter lac operator ..AGATCTCGATCCCGCGAAATTAATACGACTCACTATAGGGGAATTGTGAGCGGATAACAATTCCCCTCTAGAAA Ndel rbs **ECFP** TAATTTTGTTTAACTTTAAGAAGGAGATATACATATGGTGAGCAAGGGCGAG...681nt...ATGGACGAGCTG MetValSerLysGlyGlu...227aa...MetAspGluLeu ECFP Sacl HinDIII EcoRI Ncol BamHI Eagl Xhol TACAAGAGCTCAAGCTTGAATTCTTCGGCCGGAGGCAGCAACGGCTCGAGCGGGGTCCTCCATGGTGAGCAAG TyrLysSerSerSerLueAsnSer<u>SerAlaGlyGlySerAsnGlySerSer</u>GlyGlySerSerMetValSerLys 9-mer spacer **EYFP** His-Tag GGCGAG...681nt...ATGGACGAGCTGTACAAGGTCGAGCACCACCACCACCACCACTGAGATCCGGCTGCTA GlyGlu...227aa...MetAspGluLeuTyrLysValGluHisHisHisHisHisHisEnd T7 Terminator ACAAAGCCCGAAAGGAAGCTGAGTTGGCTGCTGCCACCGCTGAGCAATAACTAGCATAACCCCTTGGGGCCTCTA...

Figure S3. Vector pKC1

ECFP was digested with NdeI and BamHI. EYFP was digested with BamHI and SalI. pET21a was digested with NdeI and XhoI. The three fragments were ligated together to produce pKC1 (figure S3).

### The oligos used to create the IKK FRET sensors are:

IKK1F: 5'-GGTTGGAAGCTTGGAGCGTCTGCTCGACGATCGCCATGACAGCGGACTGG

IKK1R: 5'-GGTGGTCGGCCGATTCCTCATCCTTCATAGAATCCAGTCCGCTGTCATGG

IKK2F: 5'-GGTGGTCTCGAGCGGAAGCAACGGTGGAAGCGTCTGCTCGACGATCGCCATGACAGC

IKK2R: 5'-GGTGGTGGATCCAGATTCCTCATCCTTCATAGAATCCAGTCCGCTGTCATGGCGATCGTCG

C25XhoI5S: 5'-GGTGGTCTCGAGCATGCTGGAGGTCGTCGAAGCATC

C25XhoI5L: 5'-GGTGGTCTCGAGCGGAAGCAACGGTGGAAGCAACGGGTCGTCTATGCTGGAGGTCGTCGAAGC

C25HindIII5: 5'-GGTTGGAAGCTTGGGCGGTATGCTCGAGGTCGTCGAAGC

FnHindIII5: 5'- GGTTGGAAGCTTGGGCGGTATGCTCGAGGTCGTCGCTGC

FnEag13S: 5'- GGTGGTCGGCCGAGGTGCGGTAGTTGATGGAGATCG

FnEagl3L: 5'- GGTGGTCGGCCGACCCGTTGCTTCCACCGTTGCTTCCGCTAGAGGTGCGGTAGTT-

GATGGAGATCG

First, the IKK substrate sequences were assembled by extending either IKK1F and IKK1R or IKK2F and IKK2R with Klenow Fragment (3'→5' exo-) per the manufacturer's instructions (NEB). To create IKK FS C25 1S and 1L, the IKK1 fragment was digested with HindIII and EagI and ligated into the HindIII and EagI restriction sites of pKC1. To create IKK FS C25 2S and 2L, the IKK2 fragment was digested with XhoI and BamHI and ligated into the XhoI and BamHI sites of pKC1. 10C17C25 was amplified with either C25XhoI5S and Fnoligo8 or C25XhoI5L and Fnoligo8. Both PCR products were digested with XhoI and BamHI and ligated into pKC1 containing the IKK1 substrate sequence to create IKK FS C25 1S and 1L (See Figure 4b and Figure S4). 10C17C25 was next amplified with either C25HindIII and FnEagI3S or

C25HindIII and FnEagI3L. A non-selected control, Fn04, was also amplified with FnHindIII5 and FnEagI3S. All three PCR products were digested with HindIII and EagI and ligated into pKC1 containing the IKK2 subtrate sequence to create IKK FS C25 2S, 2L and control FS Fn04. All spacer sequences are listed in figure S4.

```
C25 1S CFP SSSL IKK SUBSTRATE SAGGSNGSS Fn10C17C25 GSS YFP

C25 1L CFP SSSL IKK SUBSTRATE SAGGSNGSSGSNGGSNGSS Fn10C17C25 GSS YFP

C25 2S CFP SSSLGG Fn10C17C25 SAGGSNGSSGSNGGS IKK SUBSTRATE SGSS YFP

C25 2L CFP SSSLGG Fn10C17C25 SSGSNGGSNGSSGSNGGS IKK SUBSTRATE SGSS YFP

Fn04 CFP SSSLGG Control Fn04 SAGGSNGSSGSNGGS IKK SUBSTRATE SGSS YFP
```

Figure S4. IKK FRET sensors. The underlined spacer sequences are encoded by pKC1.

#### REFERENCES

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- (3) Ja, W. W., Adhikari, A., Austin, R. J., Sprang, S. R., and Roberts, R. W. (2005) A peptide core motif for binding to heterotrimeric G protein alpha subunits. *J Biol Chem* 280, 32057-60.