Mechanism of Endogenous Regulation of the Type I Interferon Response by Suppressor of IκB Kinase ε (SIKE), a Novel Substrate of TANK-binding Kinase 1 (TBK1)*

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Background: Suppressor of IκB kinase ε (SIKE) inhibits a key innate immune effector molecule, TANK-binding kinase 1 (TBK1), through an undefined mechanism.

Results: SIKE is a TBK1 substrate.

Conclusion: SIKE controls TBK1 activity by acting as a high affinity substrate.

Significance: SIKE attenuates phosphorylation of interferon regulatory factor 3 (IRF3) by serving as an alternative, high affinity substrate for TBK1.

TANK-binding kinase 1 (TBK1) serves as a key convergence point in multiple innate immune signaling pathways. In response to receptor-mediated pathogen detection, TBK1 phosphorylation promotes production of pro-inflammatory cytokines and type I interferons. Increasingly, TBK1 dysregulation has been linked to autoimmune disorders and cancers, heightening the need to understand the regulatory controls of TBK1 activity. Here, we describe the mechanism by which suppressor of IκKe (SIKE) inhibits TBK1-mediated phosphorylation of interferon regulatory factor 3 (IRF3), which is essential to type I interferon production. Kinetic analyses showed that SIKE not only inhibits IRF3 phosphorylation but is also a high affinity TBK1 substrate. With respect to IRF3 phosphorylation, SIKE functioned as a mixed-type inhibitor ($K_{i,\text{app}} = 350 \text{ nM}$) rather than, given its status as a TBK1 substrate, as a competitive inhibitor. TBK1 phosphorylation of IRF3 and SIKE displayed negative cooperativity. Both substrates shared a similar $K_v$ value at low substrate concentrations (~50 nM) but deviated >8-fold at higher substrate concentrations (IRF3 = 3.5 μM; SIKE = 0.4 μM). TBK1-SIKE interactions were modulated by SIKE phosphorylation, clustered in the C-terminal portion of SIKE (Ser-133, -185, -187, -188, -190, and -198). These sites exhibited striking homology to the phosphorylation motif of IRF3. Mutagenic probing revealed that phosphorylation of Ser-185 controlled TBK1-SIKE interactions. Taken together, our studies demonstrate for the first time that SIKE functions as a TBK1 substrate and inhibits TBK1-mediated IRF3 phosphorylation by forming a high affinity TBK1-SIKE complex. These findings provide key insights into the endogenous control of a critical catalytic hub that is achieved not by direct repression of activity but by redirection of catalysis through substrate affinity.

To defend against pathogens, multicellular organisms mount an immune response that recognizes, sequesters, and eradicates invading infectious agents. Essential to this safeguard is the innate anti-viral response mediated by type I interferons (1–3). To initiate type I interferon production, pattern recognition receptors, such as Toll-like receptor 3 (TLR3)2 (4), retinoic acid-inducible gene I (5), melanoma differentiation factor 5 (6), or DNA-dependent activator of interferon-regulatory factors (7), recognize and respond to distinct virus-associated molecular signatures, for example double-stranded RNA (dsRNA). These receptors trigger signaling cascades that converge at the kinase, TANK-binding kinase 1 (TBK1) (8–10).

TBK1 (11), also known as NF-κB-activating kinase (12) or TRAF2-associated kinase (13), is a ubiquitously expressed serine/threonine kinase of the IκB kinase family. The canonical IκB kinases, IKKα and IKKβ, phosphorylate IκBα allowing the transcription factor, NF-κB, to induce a proinflammatory response (14). TBK1, a noncanonical IκB kinase, targets the transcription factors interferon regulatory factor 3 (IRF3) and 7 (9). In IRF3, seven Ser/Thr sites are phosphorylated near the C terminus.385SSX386SSXXX387X388X389X390 (15, 16). Of these seven phosphorylation sites, phosphorylated Ser-396 to Ser-405 induces IRF3 interaction with CBP/p300, whereas phosphorylation at residues Ser-385 and Ser-386 controls IRF3 dimerization (17–20). Modified IRF3 translocates to the nucleus where it binds to the interferon β enhancer and contributes to type I interferon production (15). In addition to IRF3/7, several other substrates have been identified that implicate TBK1 function in the insulin response (insulin receptor (21), cell growth (Akt (22)) and xenophagy (optineurin (23, 24))). Not surprisingly,

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aberrant TBK1 activity contributes to autoimmune disorders and cancers (22, 25–27).

To become active, TBK1 requires phosphorylation at Ser-172 within the kinase domain activation loop (28). IKKε (29) and TBK1, via a trans-phosphorylation event (30), provide this modification. Phosphorylated Ser-172 induces reorganization of the activation loop, making the active site competent to bind substrate (30). Alignment of TBK1 substrates revealed a weak consensus sequence for TBK1 phosphorylation consisting of a hydrophobic residue immediately preceding the modified serine that is mirrored by several hydrophobic residues lining the P + 1 site (30). Kinetic analysis of TBK1 phosphorylation of Ser-36 within the IκBα peptide (residues 19–41) revealed that TBK1 functions through a rapid equilibrium, random order mechanism (31), consistent in all IκB kinases (31–33).

To regulate TBK1 activity, endogenous inhibitors that alter TBK1 ubiquitination (A20 (34)) or mask tyrosine phosphorylation sites (SHP-2 (35)) have been characterized. In addition, a TBK1 ubiquitination (A20 (34)) or mask tyrosine phosphorylation (34) or mask tyrosine phosphorylation as a TBK1 substrate. 

**Constructs**—The human SIKE sequence was cloned from total RNA isolated from the 786O cell line using the One-step RT-PCR kit according to the manufacturer’s protocol (Qiagen). The primers incorporated a 5’ Ndel restriction enzyme site (5’ ATTATCATATGAGCAGCTGACCATCGAAGATC 3’) and a 3’ BamHI restriction enzyme site (5’ TAATAGGATCTTATTTGTGCTTGGGAAGC 3’). The vector was used to inoculate an overnight culture of Luria broth medium (RPMI 1640 supplemented with 10% low endotoxin horse serum, 20 mM l-glutamine, 100 mM HEPES, 10 mM sodium pyruvate, and 1× nonessential amino acid solution) at 37 °C in 5% CO2.

**Expression and Purification of Proteins**—For pET15b SIKE72 construct expression, the vector was transformed into chemically competent BL21-CodonPlus (DE3)-RIPL following the manufacturer’s protocol (Agilent Technologies). SIKE(1–112) and 184 (SIKE(72–184)) and the N-terminal directed mutagenesis kit following the manufacturer’s protocol (Agilent Technologies). All constructs and constructs were confirmed by DNA sequencing.

**EXPERIMENTAL PROCEDURES**

**Reagents**—Chemicals were purchased from Sigma unless otherwise specified. [γ-32P]ATP was purchased from Perkin-Elmer Life Sciences. Recombinant TBK1 was purchased from Invitrogen. Anti-phosphoserine antibody was purchased from Qiagen. Anti-FLAG-HRP was purchased from Sigma. Anti-HA antibody was purchased from Covance. Anti-full-length IRF3 and anti-actin-HRP antibodies were purchased from Santa Cruz Biotechnology. Anti-phospho-Ser-396 IRF3 antibody was purchased from Cell Signalling. Anti-rabbit-IgG-HRP and anti-mouse-IgG-HRP antibodies were purchased from Southern Biotechnology Associates, Inc.
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buffer. Bound protein was refolded on the column using a 40-CV reverse gradient of GdnHCl buffer to 100 mM sodium phosphate, pH 8.0, 500 mM NaCl, and 1 mM 2-mercaptoethanol (Buffer 1) and eluted with 5 CV of Buffer 1 plus 500 mM imidazole. Elution fractions were separated on a HiLoad 16/60 Superdex 200 column (GE Healthcare), and peak fractions were screened for target protein by SDS-PAGE analysis. CD and thermal denaturation of WT and mutant SIKE72 proteins indicated primarily helical secondary structure consistent with a predicted coiled coil domain structure and a ≥2-state unfolding curve typical of globular, oligomeric proteins (data not shown).

For GST-IRF3(173–427) expression, pGEX4T1-IRF3(173–427) was transformed into chemically competent BL21-Codon-Plus (DE3)-RIPL following the manufacturer’s protocol (Agilent Technologies). Cell culture was identical to pET15b expression prior to induction. When cell density reached A600 = 0.6, cultures were incubated on ice for 30 min, induced with 1 mM isopropyl β-D-1-thiogalactopyranoside, and allowed to grow at 16 °C for an additional 14 h. Cells were harvested by centrifugation at 7,000 × g, and the cell pellet was resuspended in Buffer A (2X PBS, 10 mM DTT, 1 mM EDTA), 5 ml of buffer/g of cell pellet. Cells were emulsified to lyse (Emulsiflex C3, Avetin) and clarified by centrifugation at 12,000 × g. The supernatant was mixed with glutathione-Sepharose 4b resin pre-equilibrated in Buffer A (GE Healthcare), 5 ml. The resin/supernatant solution was loaded into a column (Bio-Rad) and washed by gravity with 100 CV of Buffer A, 100 CV of 20 mM Tris, pH 7.5, 500 mM NaCl, 1 mM EDTA, 10 mM DTT, 10% glycerol (Buffer B) and eluted with 2–5-CV fractions of Buffer B plus 10 mM glutathione, pH 8.0. The elution fractions were concentrated in an Amicon Ultra-15 centrifugal filter (Millipore) to 6 mg/ml. GST-IRF3(173–427) was incubated with 20 units/ml thrombin (GE Healthcare) for 24 h at 4 °C. IRF3(173–427) was separated from GST using a HiLoad 16/60 Superdex 75 column (GE Healthcare) equilibrated in 20 mM HEPES, pH 7.5, 500 mM NaCl, 1 mM 2-mercaptoethanol. Peak fractions were screened for cleaved, pure IRF3(173–427) by SDS-PAGE analysis. Fractions were pooled and stored at 4 °C.

Protein concentrations were determined using the Bradford method (Bio-Rad).

In Vitro Kinase Assays—Michaelis-Menten kinetic assay reactions (50-μl volume) contained 0.1 mCi of [γ-32P]ATP, 100 μM ATP, and 0.042–20.8 μM IRF3(173–427) for IRF3-varied assays; 0.1 mCi of [γ-32P]ATP, 20 μM IRF3(173–427), and 0.48–83 μM ATP for ATP-varied assays, and 0.1 mCi of [γ-32P]ATP, 100 μM ATP, and 0.043–8.4 μM SIKE72 for SIKE-varied assays. Reactions were initiated by addition of 29.6 nM and 4.93 nM final concentration, pre-activated as above, to derive initial rates for IRF3(173–427) phosphorylation. Data were plotted as the percentage of inhibited reaction rate versus SIKE72 concentration. Errors were reported as standard deviation. To derive the Ki app parameter for SIKE72 mutants, inhibited reaction rate versus SIKE72 concentration was fit to a two-parameter rectangular hyperbola.

DNA Transfection, Immunoprecipitation, and Immunoblot Analysis—Approximately 0.5 × 10^6 cells were plated into 10-cm^2 wells and transfected with 2.5 μg of total DNA of the different expression plasmids (1:0.9:0.6 ratio of pUNO-FLAG-TBK1/pCDNA3.1/pCMV-HA-SIKE) using Lipofectamine 2000 following the standard procedure (Invitrogen). After 24 h, cells were stimulated with 50 μg/ml polyinosinic:polycytidylic acid (poly(I:C)) for 3 h, harvested, and lysed in a lysis buffer (200 μl, 0.02 M HEPES, pH 7.4, 0.15 M NaCl, 10 mM NaF, 2 mM DTT, 2 mM EGTA, 1.5 mM MgCl2, 1 mM Na3VO4, 2.7 mg/ml β-glycerocephosphate, 1 mg/ml N-ethylmaleimide, 0.5% Triton X-100, 1X Complete, EDTA-free protease inhibitor mixture (Roche Applied Science)). Lysates were cleared by centrifugation (14,000 × g for 30 min at 4 °C). Protein concentration was quantified by the Bradford method (Bio-Rad). For whole cell lysates, 50 μg of total protein per sample were boiled in sample buffer (Invitrogen), separated by SDS-PAGE (10% Tris/glycine), and transferred to nitrocellulose membrane. The membrane was blocked in 5% nonfat dry milk diluted in Tris-buffered saline (TBS) containing 0.1% Tween 20 and probed with the indicated antibodies (see supplemental Table 1). Blots were
developed with chemiluminescent reagents, ECL Plus (GE Healthcare). For immunoprecipitations, cell lysates (500 μg) were incubated with 40 μl of anti-FLAG M2 antibody affinity gel (Sigma) or anti-HA resin (Sigma) for 24 h in TBS at 4 °C. Resin was washed with TBS (3× 1 ml), and bound proteins were eluted with 100 μl of 125 ng/μl FLAG peptide (Sigma) or HA peptide (Sigma). Eluted proteins were analyzed by immunoblot as described above. Each experiment was repeated in triplicate. A list of antibody dilutions for immunoblots is given in the supplemental Methods.

Phosphopeptide Mapping—Recombinantly expressed SIKE72 (500 ng) was incubated with 4.93 nM TBK1 and 100 μM ATP in assay buffer (100 μl volume) for 24 h at 30 °C. The reaction was terminated by diluting the reaction 1:1 with 6 M GdnHCl buffer. SIKE72 was isolated from the reaction by incubation with 40 μl of nickel-nitrilotriacetic acid resin, washed with GdnHCl buffer, and bound SIKE eluted with 50 μl of GdnHCl buffer plus 0.5 M imidazole. Eluent was separated by SDS-PAGE (10% Tris/bicarbonate) or anti-HA resin (Sigma) for 24 h in TBS at 4 °C.

Sequest search algorithm against custom databases generated with the double play capability of the instrument acquiring full scan mass spectra to determine peptide molecular weights and product ion spectra to determine amino acid sequence in sequential scans. The data were analyzed by database searching using the Sequest search algorithm against custom databases generated from the SIKE72 sequence. All potential phosphopeptides were manually examined for correct identification of the modified site.

RESULTS

SIKE Acts as a Mixed Type Inhibitor of TBK1-mediated Phosphorylation of IRF3—SIKE was originally classified as a physiological suppressor of TBK1 (36). Our studies into the mechanism of SIKE’s inhibitory activity focused on the TBK1-mediated phosphorylation of IRF3. The IRF3 construct used in these studies included residues 173–427, encompassing the IRF activation domain and TBK1 phosphorylation sites (385, 386, 396, 398, 402, 404, and 405). The Michaelis-Menten plot of IRF3 as the varied substrate gave initial kinetic parameters for $K_m$ (Michaelis constant) and $V_{max}$ (maximum velocity) as 2.69 μM and 10.5 nm/min reaction, respectively (Fig. 1A and Table 1).
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SIKE72 phosphorylation increased (Fig. 2A), a synthetic dsRNA mimic mimetic (Fig. 2B).

TBK1 Phosphorylates SIKE on Six Serine Residues That Mimic the IRF3 Phosphorylation Motif—To determine the number of SIKE phosphorylation sites, we assessed TBK1-mediated phosphorylation by MALDI-TOF mass spectrometry. Over a 96-h time course, we observed a mixture of singly to triply phosphorylated SIKE72 (supplemental Fig. S2A). Under the same conditions, a single to 7 out of 7 known IRF3 phosphorylation sites were modified, whereas glyoxosomal malate dehydrogenase, not known to be phosphorylated but containing 16 preferred Ser/Thr motif sites out of 36 total Ser/Thr residues, remained unmodified (supplemental Fig. S2, B and C). To identify the positions of TBK1-mediated phosphorylation in SIKE72, we completed phosphopeptide mapping by tandem mass spectrometry. Six SIKE72 phosphoseresidues (Ser-133, Ser-185, Ser-187, Ser-188, Ser-190, and Ser-198) were identified (Fig. 3, A and B, and supplemental Fig. S3, A–F). When compared with the multiple phosphorylation sites of IRF3, SIKE72 and IRF3 share a remarkable, conserved phosphorylation motif (Fig. 3C).

SIKE Phosphorylation Status Alters Its Inhibitory Activity—To determine the relationship between the multiple SIKE phosphorylation sites and SIKE-mediated inhibition of IRF3 phosphorylation by TBK1, we probed these sites via truncated mutants or site-directed mutagenesis and assessed the effect of these SIKE mutations on TBK1-mediated IRF3 phosphorylation. The broad effect of SIKE phosphorylation was analyzed by the phosphomimetic mutant, Ser to Glu mutation, at the six

| **SIKE** | **nM** | **ATP** | **K<sub>i</sub> app** | **Slope** |
|----------|--------|---------|---------------------|-----------|
| 20.8 nm  | 133.6  | 0.042–0.42 μM | 71.7 ± 5.0 | 412 ± 5.0 | 589.7 ± 4.6 | 243.4 ± 5.3 |
| 83.3 nm  | 110.4  | 0.042–0.42 μM | 58.1 ± 8.1 | 100.4 ± 4.7 | ND | 88.9 ± 3.5 |

* K<sub>i</sub> app refers to parameter describing K<sub>i</sub> for all IRF3 phosphorylation sites assessed. K<sub>i</sub> derived from intercept<sub>intercept<sub>p-IRF3; p-SIKE</sub>dS<br/>

**SIKE construct included residues 173–427 and parameters derived from data at listed concentrations.**

1). Similarly, using ATP as the varied substrate, initial kinetic parameters for the K<sub>m</sub> and V<sub>m</sub> values of the reaction were 35.3 μM and 12.3 nm/min reaction, respectively (Fig. 1B and Table 1). To examine the effect of SIKE72 on IRF3 phosphorylation, SIKE72 was added at ~20 and ~80 nM to the IR-varied or ATP-varied (Fig. 1, C and D) assays. The primary effect of SIKE72 for IRF3-varied reactions occurred on V<sub>m</sub> (~1.8-fold decrease), indicative of a noncompetitive inhibitor (Table 1). Interestingly, the K<sub>m</sub> and V<sub>m</sub> values calculated from double-reciprocal plots of the ATP-varied reactions indicated a 2.2–3-fold change in both K<sub>m</sub> and V<sub>m</sub> values consistent with a mixed-type inhibitor (Table 1). The K<sub>i</sub> app values were calculated from the double-reciprocal plot slope and intercept parameters using Equations 1 and 2, as described under “Experimental Procedures” (Table 2).
identified sites (S6E), and phospho-knock-out mutant, Ser to Ala mutations, at the six identified sites (S6A). Subsets of phospho-knock-out mutants were created to further probe sites essential for SIKE’s inhibitory activity: S4A (S185A, S187A, S188A, and S190A), S2A (S133A and S198A), and S185A. SIKE’s phosphorylation sites cluster in the C-terminal portion of SIKE. To examine the role of the N-terminal sequence in SIKE inhibition, SIKE truncation mutants containing one (SIKE(72–184)) or retaining all phosphorylation sites (SIKE(113–207)) were constructed. The panel of constructs is summarized in Fig. 4A.

Inhibition curves for the phosphomimetic and phospho-knock-out mutants showed that phosphorylation of SIKE72 reduced SIKE’s ability to inhibit TBK1-mediated IRF3 phosphorylation, whereas unmodified SIKE72 had greatly enhanced inhibitory activity (supplemental Fig. S4A). Using four concentrations that defined SIKE’s inhibitory effect, kinase assays were completed with the full panel of mutants (Fig. 4B). Apparent inhibition constants (\(K_{i,app}\)) for each mutant were derived (Table 3). The \(K_{i,app}\) for the phosphomimetic mutant increased ~3-fold over WT SIKE72, suggesting that phosphorylation may reduce the affinity between TBK1-SIKE and/or enhance release of SIKE from TBK1. The \(K_{i,app}\) for the C-terminal 113–207 construct, retaining all phosphorylation sites, was similar to WT SIKE72. Surprisingly, the \(K_{i,app}\) for the N-terminal 72–184 construct was 196 nM, even though maximal inhibition was only ~25% of the reaction. Loss of the two peripheral phosphorylation sites, Ser-133 and Ser-198, in the S2A construct also did not alter the \(K_{i,app}\) value, whereas phospho-knock-out of the four clustered serines, Ser-185, Ser-187, Ser-188, and Ser-190, reduced the \(K_{i,app}\) parameter by 2.5-fold. Within this cluster, the \(K_{i,app}\) for the point mutation S185A was similar to the S6A mutant (Table 3).

**SIKE Phosphorylation Status Controls TBK1-SIKE Interaction in Vivo**—To investigate the effect of SIKE phosphorylation on the TBK1-SIKE interaction, we utilized co-immunoprecipitation assays of epitope-tagged TBK1 and WT or mutant FL SIKE. Prior to stimulation with poly(I:C), all HA-tagged SIKE constructs co-immunoprecipitated with FLAG-tagged TBK1 (Fig. 5A). The S6E, S185E, and 1–112 constructs showed limited interaction with TBK1 relative to input protein. Following poly(I:C) stimulation, WT-FL, 1–112, and 113–207 SIKE constructs were released from TBK1, whereas the S6A, S185A, S185E, and S6E SIKE interactions with TBK1 were unchanged from unstimulated conditions (Fig. 5A). The release of WT-FL SIKE from TBK1 correlates with increased TBK1-mediated phosphorylation of IRF3 (Fig. 5B). The reduced interaction of S6E or S185E SIKE and TBK1 has no effect on dsRNA-stimulated IRF3 phosphorylation, but the stable interaction of S6A or S185A SIKE and TBK1, irrespective of dsRNA stimulation, negates dsRNA-stimulated IRF3 phosphorylation.

**SIKE Is a TBK1 Substrate**—To establish kinetic parameters for SIKE as a TBK1 substrate, TBK1 assays with saturating ATP (100 \(\mu\)M) and SIKE72 varied from 0.043 to 8.3 \(\mu\)M were completed. The \(K_m\) and \(V_{max}\) parameters were 0.41 \(\mu\)M and 7.4 nm/min reaction, respectively (Fig. 6A and Table 1). Strikingly, the double-reciprocal plot revealed a downward concave curve, similar to the double-reciprocal plot of IRF3-varied, TBK1 assays (Fig. 6B and C, and supplemental Fig. S1C).
DISCUSSION

Multiple innate immune receptors recognize pathogen-derived ligands and initiate an immediate inflammatory response (38, 39). Several of these signaling pathways converge to activate TANK-binding kinase 1 (TBK1) (9, 22, 24, 40, 41). Although a central player in the innate immune system’s defenses that shapes the downstream innate immune response, mechanisms by which the host controls TBK1 activity are not well understood. To address this issue, we sought to define mechanistically how an endogenous inhibitor of TBK1, SIKE, blocks the TBK1-mediated anti-viral response.

The major discovery of this work is that SIKE is not only an endogenous inhibitor of TBK1-mediated phosphorylation of IRF3 but is also a TBK1 substrate. SIKE has a 6.5-fold lower $K_m$ value (0.41 $\mu$M) as compared with IRF3, with a comparable $V_{\text{max}}$ value suggesting that changes in SIKE concentration could effectively manipulate TBK1 function through altered substrate selection. Using kinetic analyses and protein interaction assays, we characterized how this novel TBK1 substrate effectively inhibits type I interferon production.

**SBK1 Displays Negative Cooperativity for Substrates IRF3 and SIKE**—In our studies, we examined the effect of SIKE on TBK1-mediated IRF3 phosphorylation using a macromolecular IRF3 substrate (residues 173–427). Previous studies on the TBK1 kinetic mechanism, using an inhibitor of $\kappa$B $\alpha$ peptide as substrate, showed that the enzyme functioned via a RERO

![FIGURE 4. Phosphorylation status modulates SIKE inhibitory activity. A, schematic diagram of SIKE constructs highlighting the six potential sites of serine phosphorylation, mutations used to probe phosphorylation sites, and truncation mutants. Serines are listed from left to right: 133, 185, 187, 188, 190, and 198. CC, coiled coil. B, WT or mutant SIKE (4–1658 nm) was added to the TBK1 assay containing 5 $\mu$M IRF3 (–$K_m$ concentration), 100 $\mu$M ATP, and 4.93 nM TBK1. Reactions were completed and analyzed as described under “Experimental Procedures.”](image-url)
mechanism (31). Our data obtained with macromolecular substrates are consistent with this mechanism but reveal evidence suggesting the existence of negative cooperativity in substrate binding or the presence of a regulatory site for protein substrate. The equilibria for this mechanism with respect to IRF3 phosphorylation are given in Fig. 7A, and are described by Dalziel’s generalized rate Equation 3 (42),

\[
\frac{e}{V_o} = \frac{\Phi_0}{\Phi_1 + \Phi_2 + \Phi_3 + \text{ATP-IRF3}} \tag{Eq. 3}
\]

where \(e\) is TBK1 concentration in the reaction; \(V_o\) is the initial velocity of the reaction, and \(\Phi\) are Dalziel coefficients that, for an RERO mechanism, pertain to the concentrations of specific complexes in the reaction mechanism (\(\Phi_1 = \text{TBK1-IRF3}, \Phi_2 = \text{TBK1-ATP}, \Phi_3 = \text{TBK1}, \text{and } \Phi_0 = \text{TBK1-ATP-IRF3}\)), labeled in Fig. 7A.

We first determined the base-line kinetic parameters for IRF3 and ATP as summarized in Table 1. The double-reciprocal plot of IRF3-varied substrate was not linear but a downward concave curve (Fig. 6C). This type of plot is indicative of one of three phenomena as follows: (i) substrate activation; (ii) nonidentical active sites, or (iii) negative cooperativity. In examining these three explanations, we assumed from both the related IKK\(\beta\) (43) and TBK1 (30, 44, 45) crystallographic structure and our own size exclusion chromatography (data not shown) that TBK1 functions as a dimeric species (required for explanations ii and iii above). Substrate activation entails a substrate-binding site (with lower affinity than the active site) that when bound leads to activation of the reaction. For nonidentical active sites, the dimeric TBK1 would have a high and low affinity active site for substrate \(a\ priori\). Finally, negative cooperativity between TBK1 active sites would occur if binding of substrate at the first active site lowered the affinity for substrate at the second active site by allosteric interactions between the subunits. Alone, the IRF3 varied assay data did not allow us to discern which mechanism is at play. Luckily, SIKE also functioned as a macromolecular TBK1 substrate. The double-reciprocal plot of the SIKE-varied assays also exhibited this characteristic downward concave curvature (Fig. 6B). As SIKE and IRF3 share no sequence homology and are distinct macromolecular substrates, the substrate activation or nonidentical active site mechanism seemed improbable as these macromolecular substrates must be similarly recognized by the kinase at a site independent of the active site or with equally reduced affinity at a nonidentical active site. Rather, negative cooperativity induced by recognition of IRF3’s and SIKE’s shared phosphorylation motif could explain the nonlinear double-reciprocal plots. Negative cooperativity requires communication between active sites. Interestingly, in IKK\(\beta\) the two active sites appear to be independent of one another with dimerization mediated by the scaffold dimerization domain. In contrast, the TBK1 structure revealed three additional dimerization interfaces between the kinase and ubiquitin-like domains of one subunit to the scaffold dimerization domain of the contralateral subunit (Fig. 5).

FIGURE 5. Phosphorylation status modulates TBK1/SIKE interaction and p-IRF3 in vivo. A, HEK293 cells were transiently transfected with FLAG-TBK1 and HA-tagged WT or mutant SIKE. Cells were stimulated with poly(I:):(50 \(\mu\)g/ml) for 3 h. FLAG-TBK1 was immunoprecipitated (IP) from lysates with \(\alpha\)-FLAG resin, separated by SDS-PAGE, transferred to nitrocellulose, and immunoblotted (IB) with \(\alpha\)-HA or \(\alpha\)-FLAG antibody. Immunoblots of whole cell lysates show expression of each construct. Blots are representative of three independent experiments. B, HEK293 cells were transiently transfected with empty vector (none) or TLR3 and WT or mutant SIKE. Cells were stimulated with poly(I:):(50 \(\mu\)g/ml) for 3 h. Immunoblots of whole cell lysates were probed for \(\alpha\)-phospho-Ser-396-IRF3, \(\alpha\)-IRF3, \(\alpha\)-HA, or \(\alpha\)-actin antibodies. Blots are representative of two independent experiments. Densitometry was calculated using ImageJ software. Each band was corrected for background scatter, and the phospho-IRF3 and IRF3 bands were normalized to their respective actin bands. Ratio of phospho-IRF3/total IRF3 is reported under corresponding bands.
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FIGURE 6. SIKE is a substrate of TBK1. A, Michaelis-Menten plot of TBK1-mediated phosphorylation of SIKE with saturating ATP (100 mM), pre-activated TBK1 (4.93 nM), and SIKE varied from 0.043 to 8.4 μM. Data were fit to a 2-parameter rectangular hyperbola (SigmaPlot). B and C, Lineweaver-Burk plots of TBK1-SIKE assays with SIKE varied from 0.043 to 8.4 μM, ATP (100 mM) (B) and TBK1-IRF3 assays with IRF3 varied from 0.042 to 20.8 μM, ATP (100 mM) (C). TBK1 was at 4.93 nM in both assays. Data for 0.042–0.42 μM (C) and 1.7–20.8 μM (●) IRF3 and 0.043–0.23 μM (○) and 0.43–8.4 μM (●) SIKE were fit to a linear polynomial equation (SigmaPlot).

occurs with near equal affinity. The ~10-fold difference in $K_{m}$ values at high substrate concentration favoring the TBK1-SIKE complex would suggest that SIKE functions primarily as a competitive inhibitor of IRF3 phosphorylation.

SIKE's Inhibitory Mechanism—For a simple RERO mechanism, an alternative substrate (as we have shown SIKE to be) should give competitive inhibition with respect to IRF3 and mixed (or noncompetitive) inhibition with respect to ATP (46). Analysis of SIKE's inhibitory effects on TBK1-mediated phosphorylation of IRF3 revealed a mixed-type inhibitor (Fig. 1 and Table 1) with respect to ATP with ~2–3-fold decreases in $K_{m}$ and $V_{max}$ values (Fig. 1D and Table 1), as expected for a RERO mechanism. Surprisingly, SIKE inhibition in IRF3-varied experiments did not exhibit a competitive inhibition pattern but rather displayed primarily as a noncompetitive inhibitor, reducing $V_{max}$ by 3-fold (Fig. 1C and Table 1). To further interpret these results, we examined SIKE inhibition of the TBK1 reaction in terms of the generalized linear rate equation for a RERO mechanism developed by Dalziel (42). In the RERO mechanism, SIKE could in theory interact with any TBK1 complex and would affect the appropriate $\Phi$ parameter. The rearranged generalized linear rate Equation 4 with ATP as the varied substrate is as follows:

$$\frac{1}{V_o} = \frac{1}{ATP} (\Phi_1 + \Phi_{12} - \Phi_{12} - \Phi_1 - \Phi_{12}) \frac{1}{IRF3} + (\Phi_0 + \Phi_1 - \Phi_{12})$$  (Eq. 4)

In Equation 4, the slope effects result from SIKE interactions with $E$ and/or $E$-IRF3, whereas intercept effects result from the interaction of SIKE with $E$-ATP and/or $E$-ATP-IRF3. From the $K_{i, app}$ values determined from the slopes and intercepts of the double-reciprocal plots, we observed that the $K_{i, app}$ derived from the intercept is ~56 nM, whereas the $K_{i, app}$ derived from the slopes is ~420 nM. This suggests that the apparent affinity of the $E$-ATP or $E$-ATP-IRF3 complexes for SIKE is greater than the affinity of $E$ alone or the $E$-IRF3 complex. Rearranging the generalized Equation 5 for IRF3 as the varied substrate gives

$$\frac{1}{V_o} = \frac{1}{IRF3} (\Phi_2 + \Phi_{12} - \Phi_{12} - \Phi_1 - \Phi_{12})$$  (Eq. 5)

Here, the slope effects can be attributed to the interaction of SIKE with $E$ and/or $E$-ATP, whereas intercept effects are attributable to SIKE interactions with $E$-IRF3 and/or $E$-ATP-IRF3. The $K_{i, app}$ values determined from slope and intercept were 111 and 98 nM, respectively, equivalent between the two grouped complexes. Because we have shown that SIKE is directly phosphorylated by TBK1 (Figs. 2C and 6A), SIKE would be expected to bind to free enzyme or enzyme-ATP complexes but not IRF3-containing complexes. The fact that SIKE alters either or both $\Phi_0$ or $\Phi_1$ suggests that these effects must arise from either allosteric interactions between active sites or the existence of a separate regulatory site for SIKE. Both of these possibilities are consistent with the observed downward curvature of the double-reciprocal plots with either SIKE or IRF3 as the varied substrate. Because IRF3 and SIKE are quite different overall structures, these studies suggest that TBK1 substrates bind with negative cooperativity between otherwise identical
sites rather than the existence of an IRF3/SIKE-binding site separate from the active sites in the dimer.

**Effect of SIKE Phosphorylation on Inhibitory Function**—Four of the six phosphorylated serines of SIKE align well with the phosphorylation sites of IRF3 (Fig. 3C) (15, 16). The strongest homology was evident at the SIKE sequence ^184^LSISSE^189^ composed of a cluster of three phosphorylation sites. When TBK1 substrates are compared, the phosphorylation pattern of TBK1 substrates has three forms as follows: singly as in Akt (22), insulin receptor (21), and optineurin (23), multiple sites but not clustered as in DDX3 (47), or multiple clustered sites as in IRF3 (15, 16), IRF7 (48, 49) and, as we show here, SIKE. In IRF3/7, these clustered phosphorylation sites, when modified, alter protein function, activating the transcription factor (17–20).

We explored the role of these clustered SIKE phosphorylation sites in the context of inhibition of TBK1-mediated IRF3 phosphorylation. When replaced with the phosphomimetic mutants, Ser to Glu, SIKE's inhibitory activity (K<sub>a</sub> app ~900 nM versus WT K<sub>a</sub> app = 350 nM) and interaction with TBK1 are greatly diminished (Figs. 4B and 5A). As the introduced negative charge grossly mimics SIKE's product form, these results are consistent with product release from an enzyme. Initial studies of SIKE held that SIKE maintained TBK1 in an inactive state prior to TBK1 activation at which point SIKE was released from the TBK1 complex. Here, we attribute the TBK1-SIKE interaction to SIKE's high affinity for the enzyme and SIKE's release to its post-translational modifications.

Similar to IRF3 where individual phosphorylation sites have been attributed to specific function, we probed the six SIKE phosphorylation sites to determine whether a subset of residues was essential to TBK1's preferential binding to SIKE over IRF3. With a complete phospho-knock-out mutant (Ser to Ala), SIKE's inhibitory activity increased 4.5-fold, consistent with retained TBK1 recognition. The S2A versus S4A mutants further defined the cluster of phosphorylation sites centered at the ^185^SXSSXS^190^ motif as essential to phosphorylation-dependent release of SIKE from TBK1, whereas the phosphorylation state of the S2A sites did not contribute to SIKE's release from TBK1. The single mutation, S185A, alone could enhance inhibition of TBK1-mediated IRF3 phosphorylation. In fact, the K<sub>a</sub> app values for the S6A (77 nM) and the single S185A (41 nM) mutants are in line with the Michaelis constant at low SIKE concentrations (55.9 nM, Fig. 6B). Additional interactions between TBK1 and SIKE or the structural context of the phosphorylation motif may also contribute to the high affinity TBK1-SIKE interaction. This is supported by the ability of SIKE(72–184) to inhibit TBK1 function and SIKE(1–112) to form an interaction with TBK1 (Figs. 4B and 5A). SIKE(72–184) retains a single phosphorylation site (Ser-133) that when mutated to Ala (S2A mutant) does not alter SIKE inhibitory activity. Therefore, inhibition by this mutant is attributed to not only Ser-133 docking to TBK1's active site, but also interactions between SIKE and TBK1 outside of the active site. Similarly, SIKE(1–112), as it contains no phosphorylation sites, must form a TBK1-SIKE interaction exclusive of the active site. These interactions independent of the active site are consistent with the noncompetitive/mixed type inhibition mediated by SIKE.

**SIKE's Role in Innate Immunity**—SIKE's original characterization as an endogenous inhibitor of the TBK1-mediated antiviral response only partially explains the SIKE function. Comparison of the pseudo-K<sub>d</sub> value for the E-ATP complex showed that SIKE is preferred as a TBK1 substrate 6.5-fold over IRF3. At equivalent concentrations, SIKE would appear to inhibit IRF3 phosphorylation. This would not be the first time a TBK1 substrate had initially been defined as an inhibitor. Optineurin, an autophagy receptor, was classified as an endogenous TBK1 inhibitor (50). Subsequent studies defined optineurin as a TBK1 substrate (23, 24). Phospho-optineurin has increased affinity for the microtubule-associated protein light chain 3, thereby promoting selective autophagy of ubiquitin-coated cytosolic bacteria and providing a mechanism by which TBK1 functions in maintaining xenophagosomes (24). If not just an endogenous regulator of TBK1-mediated phosphorylation of IRF3, what is the primary function of SIKE? Two pieces of data suggest a
SIKE Is a TBK1 Substrate

SIKE function. First, a related protein, fibroblast growth factor receptor 1 oncprotein partner 2 (51), shares 56% sequence identity with SIKE but is expressed from chromosome 12 instead of chromosome 1 and lacks all six TBK1 phosphorylation sites. Fibroblast growth factor receptor 1 oncprotein partner 2 (FGFR1OP2) associates with the actin cytoskeleton and has been linked to wound healing pathways (52). Second, we have shown by fluorescence microscopy that SIKE fused to GFP localizes to the cytosol with accumulation at actin/cytoskeleton-like structures at the cell periphery and within the lamellipodia (data not shown). We hypothesize that SIKE functions in cytoskeletal rearrangement in response to pathogen challenge through its phosphorylation by TBK1. Current studies in the laboratory are focused on testing this hypothesis.

In summary, we have revealed through kinetic analysis that SIKE is a novel, high affinity TBK1 substrate. TBK1-SIKE interactions are modulated by SIKE phosphorylation but are not limited to the typical kinase-substrate interactions within the active site. TBK1 phosphorylation of increasing IRF3 or SIKE concentrations displays negative cooperativity. The mechanism by which substrate binding is allosterically conveyed between subunits remains to be determined. With respect to IRF3 phosphorylation, SIKE effectively functioned as a mixed-type inhibitor of TBK1-mediated IRF3 phosphorylation rather than, given its status as a TBK1 substrate, as a competitive inhibitor. Together, these studies suggest that endogenous control of a critical catalytic hub is not only achieved by direct repression of activity but also by redirection of catalysis through substrate affinity. Importantly, these studies identify a novel TBK1 substrate modified in response to pathogen challenge. Further investigation into SIKE’s function in the innate immune response will uncover unique pathways and provide insight into novel mechanisms of host immune defenses.

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