Probing Kinase Activation and Substrate Specificity with an Engineered Monomeric IKK2

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ABSTRACT: Catalytic subunits of the IκB kinase (IKK), IKK1/IKKα, and IKK2/IKKβ function in vivo as dimers in association with the necessary scaffolding subunit NEMO/IKKγ. Recent X-ray crystal structures of IKK2 suggested that dimerization might be mediated by a smaller protein—protein interaction than previously thought. Here, we report that removal of a portion of the scaffold dimerization domain (SDD) of human IKK2 yields a kinase subunit that remains monomeric in solution. Expression in baculovirus-infected Sf9 insect cells and purification of this engineered monomeric human IKK2 enzyme allows for in vitro analysis of its substrate specificity and mechanism of activation. We find that the monomeric enzyme, which contains all of the amino-terminal kinase and ubiquitin-like domains as well as the more proximal portions of the SDD, functions in vitro to direct phosphorylation exclusively to residues S32 and S36 of its IκBζ substrate. Thus, the NF-κB-inducing potential of IKK2 is preserved in the engineered monomer. Furthermore, we observe that our engineered IKK2 monomer readily autophosphorylates activation loop serines 177 and 181 in trans. However, when residues that were previously observed to interfere with IKK2 trans autophosphorylation in transfected cells are mutated within the context of the monomer, the resulting S9 cell expressed and purified proteins were significantly impaired in their trans autophosphorylation activity in vitro. This study further defines the determinants of substrate specificity and provides additional evidence in support of a model in which activation via trans autophosphorylation of activation loop serines in IKK2 requires transient assembly of higher-order oligomers.

The IκB kinase (IKK) complex responds to a host of proinflammatory cell stimuli by switching to a state of high catalytic activity. The prototypical IKK complex contains three subunits: IKK1/IKKα, IKK2/IKKβ, and NEMO/IKKγ. IKK1 and -2 are closest related kinase domain-containing polypeptides, whereas NEMO is an obligate scaffolding subunit.1–3 The essential nature of NEMO in IKK activation is illustrated by the observation that NEMO−/− MEF cells are insensitive to canonical inducers of NF-κB activity, including TNF-α, LPS, and IL-1.4

Activation of IKK1 and -2 results from the phosphorylation of two activation loop serines: S176 and S180 for IKK1 and S177 and S181 for IKK2.5 Several upstream IKK kinases, including TAK1, MEKK3, and IRAK1, have been reported.6–9 No single upstream kinase, however, is required for IKK activation under all conditions of signaling and across all cell types. Furthermore, several viral gene products such as HTLV Tax and KSHV v-FLIP trigger IKK activation independent of upstream signaling by directly binding to NEMO.10–13 Additionally, it has been demonstrated in yeast and insect cells that overexpressed recombinant human IKK1 and -2 subunits can become activated without NEMO or any specific upstream kinase.14,15 Finally, the addition of free polyubiquitin chains was shown to be sufficient to activate purified IKK1 and -2 in an in vitro reconstitution of IKK activation, suggesting that activation of these kinase subunits may result from trans autophosphorylation.16

Although active IKK recognizes and phosphorylates a number of diverse cellular substrates, including FOXO3a, p65/RelA, SNAP-23, and TSC1, its best understood function is phosphorylation of the transcription factor NF-κB inhibitor protein IκBα on two N-terminal serine residues, S32 and S36.17–19 This signal-dependent phosphorylation of IκBα occurs in the cytoplasm and quickly leads to its ubiquitinylation and 26S proteasome-mediated degradation. The newly freed NF-κB then migrates to the nucleus where it elevates the expression of target genes. Mouse knockout studies clearly revealed that the IKK2 subunit is primarily responsible for this

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N-terminal phosphorylation of IκBα in response to proinflammatory NF-κB-inducing stimuli such as TNF-α or IL-1β.

Recent X-ray crystal structure analyses have revealed that IKK2 contains three distinct domains: an N-terminal kinase domain (KD) followed by a ubiquitin-like domain (ULD) and a C-terminal scaffold dimerization domain (SDD). Dimerization of IKK2 subunits is mediated entirely by the SDD. Moreover, flexibility within this elongated α-helical domain facilitates opening of the otherwise rigid IKK2 dimer (Figure 1). In their open conformation, IKK2 dimers can associate with one another to form higher-order oligomers that project their KD in a manner that supports KD−KD interactions. Introduction of mutations within surfaces that mediate these higher-order interactions disrupts IKK2 activation in transfected cells, suggesting that they facilitate KD interactions through which IKK2 can become active via activation loop trans autophosphorylation.

**EXPERIMENTAL PROCEDURES**

**IKK2 Monomer Design and Preparation of Plasmids.** Preparation of IKK2(1–420) was described previously.25

IKK2(1–420) was designed from the IKK2 X-ray crystal structure and consists of human IKK2 amino acids 11–477 joined through a Gly-Gly linker to amino acids 526–637. The IKK2(11–669)pFastBacHTb plasmid starting material was modified via a QuickChange mutagenesis protocol with the primers

IKK2Δ(477–526)Fwd:
5′-GAATTCCATGGCTTCCATGTCTGAGAGAGTGAATGGTGAGCTTAATGAATGCCAGAGCTCACCAGCTCTTGCC-3′

IKK2Δ(477–526)Rev:
5′-CCAGGAGTTCATCGTCTCTGCTCTCGCCAGAGTGAATGGTGAGCTCACCAGCTCTTGCC-3′

To replace IKK2 amino acids 478–525 with two glycine codons (underlined below), a stop codon (underlined below) was then introduced after residue 557 via mutagenesis using the following pair of primers

IKK2(637–stop)Fwd:
5′-GAAGGAGGTGTTGAGCTTAATGAATTGGAGAGTGAATGGTGAGCTCACCAGCTCTTGCC-3′

IKK2(637–stop)Rev:
5′-CGGACAACAGTCTTCATCTCTTCATCTACTATTAAAGCTCACCACCTCTTC-3′

For mutagenesis of the activation loop serines 177 and 181 to phosphomimetic glutamic acid residues, the following primers were used (mutations underlined)

IKK2EE Fwd:
5′-GAGCTGATCGGGGAGCTTTGCAACAGATGAAATTGGAGAGTGAATGGTGAGCTCACCAGCTCTTC-3′

IKK2EE Rev:
5′-GGGTCACCACAGATTATTGCTGGAAGCCTCGGAGCTCACCACCTCT-3′

Mutations were also introduced to change serines 177 and 181 to alanines to generate an inactivatable IKK2 protein as well as to mutate the aspartic acid catalytic base residue 145 to asparagine to create a kinase dead variant. The following primers were used (mutations underlined)

IKK2AA Fwd:
5′-GAGCTGATCGGGGAGCTTTGCAACAGATGAAATTGGAGAGTGAATGGTGAGCTCACCAGCTCTTC-3′
Biochemistry

**RESULTS**

**Design and Engineering of an IKK2 Monomer.** Comparative analysis of the *Xenopus* and human IKK2 X-ray crystal structures revealed that individual IKK2 dimers exhibit various degrees of opening. The *Xenopus* IKK2 structure adopts a relatively closed conformation in which two subunits of the...
dimer associate through the entire length of the SDD. A closed conformation is also observed for the IKK-related TBK1 enzyme. In TBK1, however, the KD from each protomer also contacts the SDD of its dimer-forming partner, resulting in an even more compact dimer. Two recently published crystal structures of human IKK2 reveal that the dimer can adopt significantly more open conformations in which the KD–ULD swing away from one another, suggesting that dimerization is mediated entirely through portions of the SDD that are distal to the KD.

The SDD of IKK2 and TBK1 is effectively an elongated three-helix bundle in which two parallel α-helices of roughly 115 Å in length (α3s and α6s) are connected by an equally long antiparallel stretch containing three shorter helices (α3s–α5s). The open conformations observed for the human IKK2 models suggest that the SDD mediates flexibility. Indeed, superposition of Xenopus and human IKK2 models that display differing degrees of openness reveals that the structures are all roughly equivalent within either the KD–ULD–proximal SDD portions or the distal SDD but differ to a significantly greater degree for the entire subunits (Table 1). For example, overlaying the distal portion of the SDD (a discontinuous segment consisting of amino acids 477–526 and 638–666) from each of the Xenopus IKK2 subunits upon the same portion of the human IKK2 protomers (PDB code 4E3C) results in an average root-mean-square deviation (rmsd) for Cα atoms of 1.64 Å. Superposition of the remaining portions of the same IKK2 subunits (amino acids 1–477 and 526–637) yields an average rmsd of 1.70 Å. However, when full IKK2 subunits are superimposed, the average rmsd for Cα atoms measures 2.67 Å. This suggests that the structural changes that lead to the different degrees to which IKK2 adopts its open conformation are focused at a position around Gly525 between SDD helices α3s and α4s. It also suggests that, other than this hinge point, the remaining portions of the kinase are relatively rigid.

We tested this hypothesis by designing, expressing, and purifying an IKK2 construct that lacks its distal SDD region (Figure 2). The distal portion of the SDD was removed by replacing residues 477–526 (from α2s and α3s) with a Gly-Gly linker and inserting a stop codon after residue 637 in α6s (Figure 2A,B). Recombinant baculoviruses encoding for this protein were generated and used to infect Sf9 insect cell suspension cultures. Purification by Ni affinity and size-exclusion chromatography yielded a pure protein, as judged by Coomassie-stained SDS PAGE (Figure 2C). Its elution volume on size-exclusion chromatography is consistent with the deletion protein behaving as a monomer in solution (Figure 2D). Therefore, we have dubbed this engineered enzyme IKK2mono.

Circular Dichroism Spectroscopy of IKK2mono. To confirm that deletion of the residues in question does not severely alter protein secondary structure, we carried out far-UV circular dichroism (CD) spectroscopy on IKK2EE(11–669) dimers and IKK2EEmono proteins in solution. Both spectra exhibit strong troughs at 208 and 222 nm that are characteristic of proteins with significant α-helical character (Figure 3). Analysis of the spectra using DichroWeb estimates the degree of α-helical content in IKK2EE(11–669) to be 45%. This agrees remarkably well with values calculated from Ramachandran angles of the refined crystallographic model that suggest the protein is 46% α-helical. CD recognizes IKK2EEmono as 35% α-helical. Although this is slightly lower than the 43% α-helical content calculated from a theoretical model created from simply removing residues from the IKK2EE(11–669) X-ray crystal structure, it is significantly higher than the 18% α-helix contributed by the KD alone. Moreover, the difference in measured α-helicity of IKK2 monomer and dimer is accounted for primarily by an increase in turns in the monomer, the β-sheet and disordered portions calculated from the CD spectra remain unchanged. Therefore, we conclude that the truncated SDD of our IKK2mono proteins remains predominantly α-helical after removal of three α-helical passes that mediate dimerization.

IKK2mono Retains Specificity for N-Terminal Serine Phosphorylation Sites of IkBα. It was previously demonstrated that C-terminal truncations of IKK2 that eliminate the SDD entirely result in a kinase defective in its ability to recognize S32 and S36 of IkBα in vitro. These shortened versions instead direct phosphorylation toward serine and threonine residues contained within the IkBα C-terminal PEST-like region. The Xenopus and human IKK2 X-ray crystal structures reveal that the KD, ULD, and SDD participate in extensive interdomain interactions. The introduction of point mutations to destabilize IKK2 domain–domain interfaces was shown to measurably reduce the activity of full-length IKK2 toward IkBα in vitro. 23 I KK2mono maintains each of these critical interdomain interactions.

Four different versions of the human IKK2 subunit expressed and purified from baculovirus infected Sf9 insect cell suspensions were incubated independently with five different purified IkBα substrate proteins and Mg/γ-[32P]-ATP to observe the specificity of IkBα phosphorylation in vitro (Figure 4). Both the full-length IKK2 and the IKK2(11–669) protein that was previously employed for X-ray crystal structure determination phosphorylated IkBα exclusively on S32 and S36 (Figure 4; lanes 1–11). Both of these enzymes contained the S177E and S181E activation loop mutations that render the enzyme constitutively active. To test whether this pair of activation loop Glu mutations alone is sufficient to direct substrate specificity toward the N-terminal region of IkBα, we next prepared a
showing a slight decrease, as predicted because of the removal of three transcriptional activity in response to proinflammatory signaling.

Both samples show strong α-helical signals, with the monomer showing a slight decrease, as predicted because of the removal of three largely α-helical passes at the dimer interface.

Figure 2. Engineered monomeric IKK2 (IKK2mono). (A) Schematic diagram of the domain organization of the IKK2 subunit (above) and IKK2mono (below). Coloring and abbreviations are shown as in Figure 1. Numbers correspond to the domain borders in the human IKK2 subunit, and GG refers to the diglycine linker joining SDD helices α2s and α4s. (B) Ribbon diagram model of the expected IKK2mono structure on the basis of the X-ray crystal structure of human IKK2EE(11–669). (C) Coomassie-stained 10% SDS-PAGE analysis monitoring IKK2mono purification by Ni affinity (lanes 2–6) and size-exclusion (lanes 7–9) chromatography and the final concentrated protein (lane 10). (D) Representative chromatogram of size-exclusion chromatography on IKK2mono.

Figure 3. Circular dichroism spectroscopy of purified IKK2 proteins. The mean residue molar ellipticity for IKK2EE(11–669) (gray line) is plotted as a function of wavelength. Both samples show strong α-helical signals, with the monomer showing a slight decrease, as predicted because of the removal of three largely α-helical passes at the dimer interface.

human IKK2(1–420) that lacks its entire C-terminal SDD but still bears the S177E and S181E mutations (Figure 4; lanes 12–16). Just as we observed previously when assaying a similar protein construct with an activation loop of native (S177 and S181) sequence, the enzyme lacking its SDD functioned as an efficient catalyst to phosphorylate C-terminal residues of IκBα and completely failed to recognize IκBα substrate residues S32 and S36.

Our newly engineered IKK2mono displayed identical specificity in vitro as the full-length kinase. That is, it phosphorylated IκBα exclusively at S32 and S36 (Figure 4; lanes 17–21). This observation confirms that portions of the SDD proximal to and in contact with the KD–ULD are required for the phosphorylation event that results in induction of NF-κB transcriptional activity in response to proinflammatory signaling in mammalian cells.

Solution Behavior of Engineered IKK2 Monomers. It has been shown previously that IKK2 dimerization is required for its efficient activation. For example, mutation of residues that mediate dimer contacts within the distal portion of the SDD yields an IKK2 enzyme that exhibits significantly lower in vitro catalytic activity after purification from baculovirus-infected insect cells or transfected human cells. The level of phosphorylation activity by IKK2mono in our in vitro assay was reasonably high, which is interesting considering that its activation loop serines were not mutated to constitutively activating glutamic acid residues. This suggests that, even though the IKK2mono protein purifies as a monomer in solution, it might be capable of becoming activated by trans autophosphorylation in vitro.

Our previous structure-based biochemical investigations into the mechanism of IKK2 activation led to the conclusion that low-affinity interactions between IKK2 dimers through their so-called V-shaped interface stabilizes the kinase domains in a conformation that supports KD–KD interactions and promotes activation by trans autophosphorylation (Figure 1). Under this model, the principal activating signal required immediately upstream of IKK2 activation is a simple increase in the effective concentration of IKK2. This is likely a function carried out in vivo by the scaffolding protein NEMO via a process that depends upon its interaction with polyubiquitin chains.

To observe its propensity for self-association in solution, we next determined the shape-independent molecular weight of IKK2mono at different concentrations by SEC–MALLS. We first compared IKK2mono directly with full-length IKK2 and IKK2EE(11–669), the construct that was previously employed successfully for X-ray crystallography. As suggested by size-exclusion chromatography during purification, SEC–MALLS clearly reveals that IKK2mono exists exclusively as a monomer in solution, whereas full-length IKK2 and IKK2EE(11–669) are dimers (Figure 5A). We next analyzed IKK2mono at increasing concentrations and observed that it remains a monomer even at a concentration of 187 μM (Figure 5B). These observations further support the hypothesis that IKK2 dimerization is mediated entirely by the distal portion of the SDD. Furthermore, it suggests that other additional IKK2 oligomeric binding events, such as association through the V-shaped interface that was demonstrated to promote trans autophosphorylation, are of considerably low affinity.

Activation Loop Phosphorylation by IKK2 Monomers. We previously reported that when IKK2 V-shaped interface
residues I413 and L414 were both mutated to alanine and then expressed by transfection in HEK293 cells the resulting kinase was seriously impaired in its ability to become active through activation loop phosphorylation. Similarly, double mutation of residues V229 and H232, which are positioned to mediate KD−KD interactions during trans autophosphorylation, to alanine exhibited an even more severe defect. In the case of both mutants, versions in which activation loop serines 177 and 181 were also mutated to glutamic acid displayed in vitro kinase activity equal to constitutively active versions of the wild-type IKK2 sequence.

Oligomerization of two IKK2 subunits through their V-shaped interface involves surface residues from the KD, ULD, and a proximal portion of the SDD but does not necessarily require dimerization through the distal end of the SDD. Therefore, as part of our effort to characterize oligomerization-dependent trans autophosphorylation in vitro, we employed our Sf9 insect cell expression capabilities to overexpress and purify mutant versions of IKK2 mono in which key amino acid residues were mutated. Both the IKK2 mono I413A/L414A and V229A/H232A mutant proteins express to similar levels and purify as monomers, suggesting that they are folded similarly to the IKK2 mono wild-type sequence.

It has been shown previously that overexpression of IKK2 absent NEMO in Sf9 and yeast cell cultures yields active IKK2.14,15,23 In agreement with this observation, we detect phosphorylation of activation loop serine 181 by immunoblot of overexpressed and purified IKK2 prior to the addition of ATP (Figure 6A; lanes 1−4). This zero time point S181 phosphorylation is also observed for the full-length IKK2 bearing the I413A/L414A V-shaped interface mutations, suggesting that disruption of this interface is not sufficient to impede activation loop phosphorylation when IKK2 is overexpressed in Sf9 insect cells (Figure 6A; lanes 5−8).

Interestingly, S181 phosphorylation could also be detected in the engineered monomeric IKK2 mono (Figure 6A; lanes 9−12). This indicates that dimerization is not an absolute requirement for IKK2 trans autophosphorylation. However, when both the distal SDD dimerization region and V-shaped interface are compromised, we failed to observe activation loop phosphorylation either before or after incubation with ATP (Figure 6A; lanes 12−16). This observation suggests that both dimerization through the distal SDD and oligomerization via the V-shaped interface contribute to spontaneous kinase activation when IKK2 is expressed in insect cells.

We have hypothesized that the observed S181 phosphorylation is a consequence of IKK2 trans autophosphorylation because of the relatively high kinase concentrations obtained upon Sf9 cell infection. To test if IKK2 mono is, in fact, capable of

Figure 4. IKK2 mono is specific for IkBα residues S32 and S36. In vitro kinase assays were performed with full-length human IKK2EE (lanes 1−6), IKK2EE(11−669) (lanes 7−11), IKK2EE(1−420) (lanes 12−16), and IKK2 mono (lanes 17−21) against both native sequence or S32A/S36A (AA) GST−IkBα(1−54) substrates as well as native, S32A/S36A, or C-terminal S283E/S288E/T291E/T296E/T299E (E6) mutant full-length IkBα substrates. Two-hundred nanograms of kinase was used in each assay except for the IKK2EE(1−420) reactions, which contained 400 ng of enzyme each. Substrate specificity is revealed by autoradiography (top panels) and Coomassie-stained SDS-PAGE substrate loading controls (bottom panels).

Figure 5. IKK2 oligomerization state in solution. (A) Full-length IKK2 (black line) and IKK2EE(11−669) (dark gray) are both dimers in solution, as revealed by multiangle laser light scattering following size-exclusion chromatography (SEC−MALLS). IKK2 mono (light gray) is a monomer. (B) Increasing the IKK2 mono concentration to as high as 187 μM does not significantly affect its profile as a monomer in solution.

residues I413 and L414 were both mutated to alanine and then expressed by transfection in HEK293 cells the resulting kinase was seriously impaired in its ability to become active through activation loop phosphorylation. Similarly, double mutation of residues V229 and H232, which are positioned to mediate KD−KD interactions during trans autophosphorylation, to alanine exhibited an even more severe defect. In the case of both mutants, versions in which activation loop serines 177 and 181 were also mutated to glutamic acid displayed in vitro kinase activity equal to constitutively active versions of the wild-type IKK2 sequence.23
catalyzing phosphorylation of IKK2 activation loop in trans, serines 177 and 181 of the IKK2mono were mutated to glutamic acid residues to produce a constitutively active version (referred to as IKK2EEmono). In vitro kinase reactions were then carried out using a catalytically inactive version of the full-length IKK2, in which the catalytic D145 residue was mutated to asparagine (IKK2D145N), as substrate and probed by immunoblot for S181 phosphorylation (Figure 6B). The crystallized IKK2EE(11-669) and, albeit to a lesser extent, the IKK2EEmono enzymes both were found to be capable of phosphorylating S181 of full-length IKK2D145N.

Activation of IκB Kinase Activity in Monomeric IKK2. We next assessed whether oligomerization-dependent trans autophosphorylation was sufficient to activate IKK2 independent of its ability to form dimers. Relative to the engineered IKK2mono enzyme of native sequence, purified IKK2mono enzymes that harbored mutations at their V-shaped interface residues I413 and L414 were both mutated to alanine fail to be recognized by the antibody (lanes 13–16). Western with anti-His antibody confirms the level of protein in each reaction (bottom). (B) Catalytically inactive version of full-length IKK2 in which catalytic base D145 is mutated to Asn permits direct detection of activation loop trans autophosphorylation. Western blot reveals that constitutively active versions of both dimeric and monomeric IKK2 bearing the S177E/S181E mutations (EE) phosphorylate the IKK2D145N substrate in vitro (lanes 21 and 22).

Figure 6. Activation loop phosphorylation activity of IKK2 dimers and monomers. (A) Western blot analysis with an anti-phospho-S181 antibody on recombinant human IKK2 proteins purified from baculovirus-infected Sf9 insect cells before and after treatment with Mg/ATP (top). Only IKK2mono in which V-shaped interface residues I413 and L414 are both mutated to alanine fails to be recognized by the antibody (lanes 13–16). Western with anti-His antibody confirms the level of protein in each reaction (bottom). (B) Catalytically inactive version of full-length IKK2 in which catalytic base D145 is mutated to Asn permits direct detection of activation loop trans autophosphorylation. Western blot reveals that constitutively active versions of both dimeric and monomeric IKK2 bearing the S177E/S181E mutations (EE) phosphorylate the IKK2D145N substrate in vitro (lanes 21 and 22).

Figure 7. Role of trans autophosphorylation in activation of IKK2mono. (A) Autoradiography of in vitro kinase assays with GST–IκBα(1–54) as substrate and various versions of IKK2mono as enzymes (top). Both mutation of the V-shaped (I413A/L414A) and KD–KD (V229A/H232A) interfaces yields IKK2mono proteins that almost completely lack kinase activity toward GST–IκBα(1–54) substrate (lanes 1–3). Conversion of both activation loop residues S177 and S181 to phosphomimetic glutamic acids (EE) results in IKK2mono enzymes that display significantly higher kinase activity. Coomassie-stained SDS PAGE substrate and anti-His western blot enzyme loading controls are shown below. (B) Western blot with anti-phospho-S181 antibody reveals that only native sequence IKK2mono is capable of activation loop autophosphorylation (compare lanes 6, 8, and 9; above). Anti-His western blot loading control (bottom).

results from their inability to support activation via trans autophosphorylation.

To test this hypothesis further, we next used immunodetection to probe the phosphorylation state of activation loop S181 in native sequence and interfacial mutant IKK2mono enzymes after incubation with Mg and ATP. Only IKK2mono containing the native sequence activation segment (S177 and S181) could be recognized by antibodies specific for IKK2 phosphoserine-181 (Figure 7B). As controls, we also tested the IKK2mono proteins that contained activating S177E/S181E activation loop mutations. As expected, these proteins failed to react with the anti-IKK2 pSer181 antibody,
Oligomerization-Dependent Trans Autophosphorylation of IKK2. Our in vitro experiments with the engineered monomeric IKK2 enzyme reveal that its ability to become activated via trans autophosphorylation on its activation loop serines is impaired. Moreover, although after activation it retains its specificity for S32 and S36 of IκBα, IKK2\textsuperscript{mono} displays measurably lower catalytic activity toward its namesake substrate. These observations support many previously reported studies that have concluded that IKK2 dimerization plays an integral role in its activation and consequently its ability to function as an IκB kinase.\textsuperscript{15,22,32} However, our observations also suggest that IKK2\textsuperscript{mono} harbors all of the residues necessary to mediate activation and substrate specificity.

To test the hypothesis that the V-shaped and KD–KD oligomerization interfaces observed in our X-ray crystal structure are sufficient to support IKK2 activation via trans autophosphorylation, we performed a time-course experiment in which the ability of various IKK2 dimers and monomers to phosphorylate S181 of the inactive IKK2D145N mutant was monitored by western blot (Figure 8). All of the active kinases in this experiment harbored the activating S177E/S181E mutations. At a concentration of 150 nM enzyme/300nM substrate, we observed that the dimeric IKK2EE(11–669) could turn over product, whereas various IKK2\textsuperscript{mono} mutant proteins did not display activity (Figure 8; lanes 1–8). After increasing the enzyme concentration to 300 nM, however, significant phosphorylation of the inactive IKK2D145N substrate could be detected (Figure 8; lanes 9–16). The IKK2\textsuperscript{mono} 1413A/L414A mutant that harbors mutations targeting the V-shaped interface showed significantly less activity, although activity could be detected at an enzyme concentration of 450 nM (Figure 8; lanes 17–24). Mutation of the KD–KD residues V229 and H232 yielded an IKK2\textsuperscript{mono} enzyme that failed to produce detectable amounts of S181 phosphorylation at all of the concentrations tested. This experiment supports the hypothesis that activation loop phosphorylation in trans is mediated by IKK2 oligomerization interfaces in addition to dimerization.

![Figure 8. Activation loop trans autophosphorylation by mutant IKK2\textsuperscript{mono} enzymes can be partially recovered at high concentration. The activation is monitored via western blot with anti-phospho-S181 antibody. Enzyme concentrations are given; IKK2D145N was 300 nM in each reaction. Dimeric and constitutively active IKK2EE(11–669) readily phosphorylates activation loop S181 (lanes 1 and 2). IKK2EE\textsuperscript{mono} also phosphorylates the inactive IKK2D145N substrate, although a higher concentration of enzyme is required to observe comparable levels of phosphorylation (compare lanes 3 and 4, 11 and 12, and 19 and 20). Mutation of V-shaped interface residues I413 and L414 to alanine yields an IKK2EE\textsuperscript{mono} enzyme that is even more defective in trans autophosphorylation of IKK2D145N in vitro. Mutation of KD–KD interaction residues V229 and H232 to alanine yields an IKK2EE\textsuperscript{mono} enzyme that fails to trans autophosphorylate even when provided in molar excess to the IKK2D145N substrate protein.](image)

**DISCUSSION**

Protein kinases share a conserved domain structure and catalytic mechanism for phosphoryl transfer from ATP to protein substrate.\textsuperscript{33} What distinguishes one protein kinase from another, therefore, is their ability to act upon select substrates and their mechanisms for regulating catalytic activity.\textsuperscript{34,35} IKK is unique on both fronts. IKK was first cloned biochemically because it was the only kinase activity detected in TNF-α-treated HeLa cells that could specifically phosphorylate serines 32 and 36 of the NF-κB inhibitor protein IκBα.\textsuperscript{17,16,36} Despite numerous studies aimed at uncovering a basis for this substrate specificity, it is still not perfectly clear how IKK only facilitates this chemistry. The field of regulation of IKK activity has received significantly more attention, with literally thousands of published studies. These efforts have led to many seminal discoveries, including the fundamental role of ubiquitin as an activating component of cell signaling complexes.\textsuperscript{37,38} Nevertheless, basic questions surrounding the regulatory control of IKK persist. Among these, does IKK activation require any upstream kinase? How does interaction with polyubiquitin promote kinase activity? Are the extremely rapid kinetics of IKK activation observed in cells suggestive of a relatively simple activation mechanism?

We have previously shown that a version of IKK2 containing only the amino-terminal KD and ULD recognizes and phosphorylates its IκBα substrate in vitro. However, this IKK2(1–420) fragment directs its activity toward serine and threonine residues contained within the C-terminal PEST-like region of IκBα and fails to recognize S32 and S36 altogether.\textsuperscript{25} This surprising observation could be explained by one of three different factors. First, it is possible that the activation loop of IKK2(1–420) fails to become phosphorylated in the absence of its C-terminal SDD, and in its unphosphorylated state, the kinase displays altered specificity toward IκBα. A second possibility arises from the fact that the shortened IKK2(1–420) is a monomer in solution. Therefore, it is possible that dimerization through the SDD somehow directs specificity toward IκBα. The third possibility is that elements from the SDD itself are directly involved in docking and/or presenting the IκBα substrate such that S32 and S36 can be targeted. We tested a version of the IKK2(1–420) construct in which activation loop residues S177 and S181 are mutated to phosphomimetic glutamic acid residues and found that this enzyme displays the same specificity toward the C-terminal residues of IκBα, effectively disproving the hypothesis.
that activation loop phosphorylation status dictates specificity (Figure 4; lanes 17–21).

Recent success in X-ray crystallography of nearly full-length IKK2 homodimers has revealed the domain organization and arrangement of individual subunits in the functioning dimer. The observation that these IKK2 homodimers can exhibit varying degrees of opening has allowed identification of the dimerization interface within the portion of the SDD distal to the KD–ULD. In the present study, we have engineered a minimal IKK2 that lacks the distal portion of the SDD and consequently purifies as a stable monomer. This IKK2monο protein displays ideal solution behavior. Furthermore, kinase assays reveal that IKK2monο faithfully reconstitutes S32 and S36 phosphorylation of IκBα in vitro. We conclude that elements within the proximal portion of the SDD are necessary for directing specificity of IKK2 toward IκBα S32 and S36, independent of the dimerization status of the enzyme. The IKK2 X-ray crystal structures reveal that this portion of the SDD contacts both the KD and ULD, and mutations introduced to disrupt these interdomain interactions were previously shown to interfere with catalytic activity. It is likely that the proximal SDD functions either through direct contact with the IκBα substrate and/or through allosteric action on the KD to direct the specificity of IKK2 toward IκBα residues S32 and S36. In support of this hypothesis, Liu et al., when analyzing their recently published 2.8 Å X-ray crystal structure of an asymmetric dimer of human IKK2, identified a possible substrate peptide binding pocket within a crevice created by the KD and proximal SDD. Future studies aimed at understanding how this domain arrangement affects substrate specificity and its potential as a target for drug development are underway.

We recently published experimental data, based on our own X-ray crystal structure of active human IKK2, in support of a regulatory model through which raising the local concentration of the enzyme promotes its spontaneous activation via oligomerization-dependent trans autophosphorylation. Central to this hypothesis are the observations involved between IKK2 dimers in the crystal. The open IKK2 dimer conformation permits interaction between neighboring dimers through a V-shaped interface involving portions of the KD, ULD, and proximal SDD. This interaction, in turn, stabilizes IKK2 in a conformation that can support KD–KD interactions and presumably trans autophosphorylation (Figure 1). As IKK2 dimerization relies exclusively on the distal SDD, generation of the stable, catalytically active IKK2monο enzyme has enabled further validation of this IKK2 activation mechanism.

Dimerization of IKK2 has been shown to be an absolute requirement for its activity in cells. What is not clear from these studies, however, is whether blocking dimerization abolishes the potential for IKK2 to become active or whether it simply decreases the effective IKK2 concentration and consequently the probability of trans autophosphorylation. The involvement of V-shaped and KD–KD interfaces in IKK2 activation suggests that IKK2monο, which maintains these regions intact, should support trans autophosphorylation in vitro. In this study, we observe that this is, in fact, the case. We observe that the IKK2monο activation loop becomes phosphorylated spontaneously when expressed and purified from baculovirus-infected S9 insect cell suspensions. This is also true of dimeric IKK2 enzymes, suggesting that either the infected S9 cell environment or the level of IKK2 overexpression is sufficient to support its activation via trans autophosphorylation. Strikingly, when residues that mediate the V-shaped or KD–KD interfaces are mutated, there is a complete loss of activation loop phosphorylation signal and consequently minimal activity toward IκBα. This loss in IKK2 trans autophosphorylation potential can then be partially recovered by increasing the concentration of the mutated IKK2 monomers.

Taken together, these experiments with an engineered monomeric IKK2 enzyme support an IKK2 activation mechanism in which the principal upstream event is to effectivly increase the local concentration of IKK2 dimers. IKK2 dimers can then rapidly associate through V-shaped and KD–KD interfaces and become active via trans autophosphorylation. Under standard activation conditions, polyubiquitin chains induce IKK2 proximity through their noncovalent interaction with the NEMO subunit of IKK. However, this regulated process can be circumvented by other cellular events that drive IKK2 oligomerization.

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**Notes**

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**ABBREVIATIONS USED**

CD, circular dichroism; IκB, inhibitor of NF-κB; IKK, IκB kinase; HTLV, human T-lymphotropic virus; IL-1, interleukin-1; KSHV, Kaposi’s sarcoma-associated herpesvirus; LPS, lipopolysaccharide; NF-κB, nuclear factor of κ light chain gene enhancer in B cells; SEC–MALLS, size-exclusion chromatography–multiangle laser light scattering; TNF-α, tumor necrosis factor α

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