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Crystal structure of inhibitor of κ B kinase β (IKK β)

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Abstract

Inhibitor of κ B (I κ B) kinase (IKK) phosphorylates I κ B proteins leading to their degradation and liberation of nuclear factor κ B (NF- κ B) for gene transcription. Here we report the crystal structure of IKK β in complex with an inhibitor at 3.6 Å resolution. The structure reveals a tri-modular architecture with the kinase domain (KD), a ubiquitin-like domain (ULD) and an elongated, α -helical scaffold/dimerization domain (SDD). Surprisingly, the predicted leucine zipper and helix-loop-helix motifs do not form these structures but are part of SDD. The ULD and SDD mediate a critical interaction with I κ B α that restricts substrate specificity, and the ULD is also required for catalytic activity. The SDD mediates IKK β dimerization, but dimerization *per se* is not important for maintaining IKK β activity, and instead is required for IKK β activation. Other IKK family members IKK α , TBK1 and IKKi may share the similar tri-modular architecture and function.

NF- κ B transcription factors are master regulators of inflammatory, immune and apoptotic responses 1,2. In the canonical pathway, NF- κ B dimers are held in the cytoplasm via binding to I κ B proteins, which mask their nuclear localization signals. When cells are stimulated by NF- κ B inducers, I κ Bs are phosphorylated by the Ser/Thr-specific IKK, a modification that triggers their Lys48-linked polyubiquitination and subsequent proteasomal degradation 3. Freed NF- κ B dimers enter the nucleus to regulate gene transcription 2. In the

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Author Contributions

G.X. cloned, expressed, purified, crystallized and determined the crystal structure of xIKK β and performed Km determination experiments. Y.C.L. cloned, expressed, purified and crystallized hIKK β and performed pulldown experiments and kinase assays using phospho-I κ B α antibody. Q.L. expressed the hIKK β mutants in insect cells. G.N. and X.W. vperformed transfection, immunoprecipitation and kinase assays and M.K. supervised these experiments. H.W. supervised the project. G.X and H.W. made the figures and wrote the manuscript.

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non-canonical pathway, activated IKK phosphorylates the I κ B-like domain in the NF- κ B family member p100/NF- κ B2 3. NF- κ B signaling pathways are associated with a vast number of human diseases including inflammation and cancer, which renders IKK a potentially important therapeutic target 4 (www-nf-kb.org).

IKK was originally purified from HeLa cells as a multi-protein complex that contains the kinase subunits IKK α and/or IKK β , and the regulatory protein NEMO (also known as IKK γ or FIP-3) 5–11. IKK α and IKK β both contain an N-terminal kinase domain (KD), predicted leucine zipper (LZ) and helix-loop-helix (HLH) domains, and a C-terminal NEMO binding domain (NBD) (Fig. 1a). IKK β appears to have an additional ubiquitin-like domain (ULD) following the KD, which is not predicted in the corresponding region of IKK α 3. IKK-related kinases TBK1/NAK and IKKi/IKK ϵ appear to share a similar domain organization 12. While IKK β mediates activation of the canonical NF- κ B pathway in response to pro-inflammatory stimuli, IKK α plays an indispensable role in non-canonical NF- κ B signaling by phosphorylating p100/NF- κ B2. Protein kinase assays suggested that IKK β accounts for nearly all of the catalytic activity of the IKK holoenzyme towards I κ B α 3,13.

The activation loop in both IKK α and IKK β KDs contains the MEK consensus motif SxxxS (S177 and S181 in human IKK β) 6–8,10. Some MEK kinase family members, such as TGF- β -activated kinase 1 (TAK1) and NF- κ B-inducing kinase (NIK), were shown to phosphorylate IKKs 3,14,15. IKK α and IKK β can also undergo autophosphorylation and activation as a result of overexpression or signal dependent NEMO clustering 10,16. Ala substitutions of the activation loop Ser residues prevent IKK activation whereas the phosphomimic, double Glu mutation S177E/S181E (EE) of IKK β renders it constitutively active 7,13.

Tri-modular arrangement of IKK β

We determined the crystal structure of *Xenopus laevis* IKK β (xIKK β) EE (residues 4–675) (Fig. 1a) in complex with either inhibitor Cmpd1 or Cmpd2 (Supplementary Fig. 1) at 4.0 and 3.6 Å resolutions in I4 $_1$ 22 and P1 space groups, respectively (Supplementary Table 1 and 2, Supplementary Fig. 2). Eight IKK β molecules in P1 and the one molecule in I4 $_1$ 22 are highly similar to each other (Supplementary Fig. 3, Supplementary Table 3) and show conserved dimerization (see below). Structural description will use the first dimer (chains A and B) in P1. The hIKK β and xIKK β sequences share 74% identity with no gaps within the region of the structure; residue numbers designated for xIKK β are also true for hIKK β .

The IKK β dimer structure resembles a pair of shears and has the overall dimensions of approximately 100Å x 130 Å x 60 Å (Fig. 1b, 1c). It contains KD (16–307), ULD (310–394), and a highly elongated domain we here refer to as the scaffold/dimerization domain (SDD, 410–666) (Fig. 1a, Supplementary Fig. 4). While KD and ULD form the handle of the shears, SDD is the blade. Both KD and ULD intimately associate with SDD, suggesting that KD does not function independently. Instead, the structure indicates that IKK β is an integral tri-modular unit.

IKK β KD in complex with either inhibitor Cmpd1 or Cmpd2 exhibits the typical bilobal kinase fold 17. Although sharing only 21.1% sequence identity with human ubiquitin, ULD

of IKK β (310–394) indeed has the ubiquitin fold (Fig. 1d). A major difference is the presence of an eight-residue insertion (residues 373–380) that forms part of the loop connecting $\beta 4'$ and $\beta 5'$ in ULD, which is mostly disordered. The hydrophobic surface patch of ubiquitin 18, which is often recognized by ubiquitin binding proteins, is conserved in ULD, with residues V318, L354 and L389 equivalent to L8, I44 and V70 of ubiquitin (Supplementary Fig. 5).

SDD comprises six helices ($\alpha 1s$ – $\alpha 6s$), among which $\alpha 2s$ and $\alpha 6s$ each contain 70 and 77 residues, respectively. They twist together with a stretch of three shorter helices between them to fold as an elongated structural ensemble. The KD C-lobe sits on the N-terminal end of SDD while ULD binds close to the middle of SDD. Surprisingly, formerly designated LZ (458–485) and HLH domains (605–644) 7,8,10 do not exist as such in the structure and are both parts of SDD (Fig. 1e). Most of the hydrophobic residues in the predicted LZ point inward and are not available for mediating dimerization as previously proposed.

Structure of inhibitor bound IKK β KD

The inhibitor binds to IKK β KD at the hinge loop connecting N- and C-lobes, a region that recognizes the adenine in ATP 19,20 and often used for inhibitor binding 21–23 (Fig. 2a, Supplementary Fig. 6). The KD conformation is incompatible with that of an active kinase 17,24,25 (Fig. 2b, 2c). The activation segment, which begins from D166-L167-G168 as the conserved DFG triad and extends to A190-P191-E192 in the conserved APE motif, is fully ordered, including phosphomimic residues E177 and E181 (Fig. 2b). However, the C-terminal anchor of the activation segment, including the APE motif itself, is completely out of place so that essential interactions with the catalytic center are lost (Fig. 2c). Gross conformation of the N-terminal anchor of the activation segment is preserved with $\beta 9$ paired with the $\beta 6$ strand that precedes the catalytic loop. Part of the activation loop (175–177) contains an additional β strand ($\beta 10$) that inserts in between the lobes to form a three-stranded β sheet with $\beta 9$ and $\beta 6$. The αC helix is tilted up and outward (Fig. 2c) to make room for $\beta 10$, which also disrupts its productive interactions with the DFG motif in active kinase structures.

Interactions among KD, ULD and SDD

KD, ULD and SDD interact mutually, with the ULD: SDD interaction being most extensive, followed by the KD: SDD and the KD: ULD interactions. ULD binds close to the middle of SDD, contacting helices $\alpha 2s$ and $\alpha 6s$ (Fig. 3a). The interaction buries ~ 700 Å^2 surface area per partner. There are substantial hydrophobic contributions, supplemented by electrostatic interactions. Residues M315, M317, L354, I387, L389 and F390 on one side of ULD, pack against L612, Y609, L447 and main chain of $\alpha 2s$ of SDD to form the central hydrophobic core of the interface. This hydrophobic patch of ULD is immediately adjacent to and overlaps with the conserved hydrophobic patch in the ubiquitin fold. Electrostatic interactions are observed between E352 and K619 and between K394 and D610. Additional interfacial residues include S319 and S357 of ULD and T453, Q449, and Y609 of SDD.

Consistent with an important role of ULD in interacting with SDD, mutations in residues that are not at the interface, P347A, L361A and Q351A, had minimal effects on NF- κ B

activation 26. In contrast, mutation in a residue within a ULD surface loop that contacts the SDD, G358A (Fig. 3a), affected IKK β -induced NF- κ B activity 26. It was also shown that L353 is absolutely required for IKK β activity 26; however, L353 is buried in the ULD core and the L353A mutation may have disrupted IKK β structural integrity. Double substitutions of human IKK α and IKK β that are equivalent to I608R/Y609P of SDD of α IKK β did not affect their interaction with WT IKK β , but negatively impacted kinase activity 10; I608 is buried in the SDD core and Y609 directly interacts with ULD (Fig. 3a).

Similar to ULD, KD also makes contacts with the N-terminal end of SDD (Fig. 3b), burying a surface area of $\sim 650 \text{ \AA}^2$ from each interface. The binding forces are mainly van der Waals in nature. Limited hydrophobic interactions are observed, between residues A252 and V253 of KD and Y423 of SDD, and between F111 of KD and the hydrophobic portions of R572, R575 and E576 of α 5s of SDD. KD is linked to ULD through a two amino acid linker between α I of KD and β 1' of ULD (Fig. 3c), burying only $\sim 350 \text{ \AA}^2$ surface area. Side chain contacts between L311 of ULD and I268 of KD are observed, which together form the small hydrophobic patch at the KD-ULD junction consisting also of L269 and I306 of KD and L309 of the linker. Together with the ionic interaction between D373 of ULD and R123 of α E of KD, these interactions may confer rigidity to the KD-ULD junction despite the small buried surface area.

Structural comparison with other kinase structures revealed a similarity of the locations of SDD and ULD to several known docking sites for substrates and regulatory proteins 27. In the crystal structure of the Ser/Thr kinase SRPK1 in complex with a docking motif in its substrate ASF/SF2 27, the peptide motif interacts with the kinase at the location of SDD (Fig. 3d). In the structure of the TAK1 kinase domain fused with the C-terminal region of its binding protein TAB1 28, TAB1 interacts with the C-lobe of the kinase at a position analogous to both SDD and ULD, presumably to activate the kinase (Fig. 3d).

ULD-SDD binds I κ B α C-terminal region

Previous studies have suggested that ULD in TBK1 and IKKi is involved in substrate recognition because its deletion impaired activity of the respective kinases and a GST-ULD fusion protein interacted with the specific substrates IRF3 or IRF7 29. Because ULD deletion in IKK β also abolished its activity 26, we hypothesized that the ULD may recognize its specific substrate I κ B α . However, we were surprised to find that GST-I κ B α (1–317 and 54–317) pulled down only a minute amount of ULD (Fig. 4a, lanes 9 and 13), while GST alone did not pull down any (Fig. 4a, lane 4), suggesting that the interaction of I κ B α with ULD is specific, but very weak. In contrast, GST-I κ B α robustly pulled down full-length IKK β or IKK β lacking C-terminal NBD (Fig. 4a, lanes 15, 16).

I κ B α has an N-terminal region (1–54) that contains cognate phosphoacceptor sites at S32 and S36, followed by a C-terminal region (55–317) that contains multiple ankyrin repeats and the PEST region 30,31. Strikingly, the N-terminal region of I κ B α did not pull down any IKK β constructs (Fig. 4a, lanes 5–8), while the C-terminal region of I κ B α interacted robustly with full-length IKK β as well as its ULD-SDD region (Fig. 4a, lanes 9–12), and very weakly with ULD alone (Fig. 4a, lane 9). Further mapping on I κ B α showed that both

ankyrin repeats (1–282, or 54–282) and the PEST region (282–317) interacted with IKK β ULD-SDD (Fig. 4b, lanes 4, 5, 6). The extent of pulldown suggests that the PEST region contributes more to IKK β interaction than the ankyrin repeats. Despite trying multiple constructs, we could not obtain soluble protein for IKK β SDD to test its interaction with I κ B α . However, the pull down data suggest a dominant role of SDD in I κ B α interaction. In any case, it is clear that the mutual interaction between IKK β and I κ B α is mediated by their ULD-SDD and C-terminal domains, respectively.

ULD-SDD limits specificity and ULD aids catalysis

The specific interaction between ULD-SDD of IKK β and I κ B α suggests that the K_m of phosphorylation by IKK β might be lower for full length I κ B α than for its N-terminal region (1–54) alone. We performed kinase kinetics analysis of IKK β EE against the two different substrates. Surprisingly, the measured apparent K_m values were 11.4 μ M and 13.7 μ M for full-length and the N-terminal region of I κ B α , respectively (Fig. 4c, d), suggesting that binding at SDD, an exosite, does not alter the K_m of the reaction dramatically. This could be due to opposing effects of the SDD: I κ B α interaction, which increases substrate interaction but slows down product dissociation. The relative V_{max} values were 502.4 and 250.2, respectively, suggesting that the SDD: I κ B α interaction moderately enhances catalysis.

Casein kinase 2 (CK2) phosphorylates the C-terminal PEST region of I κ B α around residues 283–299 30,31. To determine whether the SDD: I κ B α interaction restricts substrate specificity, we compared the kinase activity of purified IKK β EE proteins against either I κ B α or its S32A/S36A mutant (AA) using [γ - 32 P]ATP (Fig. 4e). Although the KD-ULD (1–400) construct gave rise to a small amount of protein, it produced robust phosphorylation on WT I κ B α , comparable to full-length IKK β (1–756), suggesting that it is catalytically competent. Remarkably, KD-ULD effectively phosphorylated the AA mutant which in contrast was a poor substrate for full-length IKK β . The C-terminal PEST region appeared to harbor the major sites of phosphorylation by KD-ULD because a construct lacking PEST (1–282) was not significantly phosphorylated by KD-ULD but was phosphorylated by full-length IKK β (Fig. 4e). In addition, when I κ B α phosphorylation was detected by antibody against I κ B α phosphorylated at S32/S36, only very weak phosphorylation was seen for the KD-ULD construct in comparison with full-length IKK β (Fig. 4f). These experiments suggest that while the KD-ULD construct is active, it possesses an altered substrate specificity, causing preferential phosphorylation of the C-terminal PEST sequence, consistent with a previous observation 32. Hence, ULD-SDD appears to specifically position I κ B α such that only its N-terminal region is accessible to the IKK β catalytic pocket (Fig. 4g).

We expressed three KD constructs, 1–301, 1–310 and 1–360, in insect cells and only obtained small amount of protein for the 1–360 construct. Kinase assay showed that IKK β EE (1–360) exhibited very low residual activity against I κ B α or its AA mutant (Fig. 4e), suggesting that KD activity is compromised in the absence of ULD. Without an isolated KD structure, we cannot deduce the molecular mechanism by which ULD acts. However, in analogy to TAK1 activation by TAB1 28 (Fig. 3d), it may be speculated that this KD-ULD interaction allosterically potentiates kinase activity. Further kinase assay using antibody

against I κ B α phosphorylated at S32/S36 showed no detectable activity of KD alone (Fig. 4f), confirming that the low residual activity may also be directed towards the C-terminal PEST region, similar to KD-ULD. Therefore, while ULD-SDD is critical for IKK β specificity, ULD is required for its catalytic activity.

SDD mediates IKK β dimerization

Full-length hIKK β (1–756) and its KD-ULD-SDD region (1–678) are dimers in solution as determined by gel filtration chromatography (Fig. 5a). In the P1 and I4₁₂₂ crystals, two conserved dimerization interfaces exist, one mediated by SDD and the other mediated by KD. Because hIKK β (1–643) that truncates into SDD is a monomer in solution (Fig. 5a), SDD-mediated dimerization (Fig. 5b) is likely of greater importance than KD-mediated dimerization.

SDD in an IKK β dimer does not form extensive interactions along the entire length dimension of the helical bundle. Rather, interactions are mostly localized to the end of the bundle (Fig. 5b), distal from KD and ULD and burying $\sim 1,000 \text{ \AA}^2$ surface area from each monomer. The interaction is mostly hydrophobic. Residues that contribute significantly to dimerization include Q478, K482, F485, I492, K496, I505, Q651, L654, W655, L658, and I660 with residues L654, W655 and L658 from helix α_6 s burying the most surface area (Fig. 5b). This dimerization interface is entirely unexpected as it was predicted earlier that LZ mediates IKK β dimerization. The structure now explains that failed dimerization of the LZ-defective L462S/L469S mutant of IKK α 10 is due to the structural role of L469, whose equivalent residue M472 of IKK β is buried in the SDD core. Superposition of four IKK β dimers in P1 and the one IKK β dimer in I4₁₂₂ shows that IKK β dimers are conserved (Supplementary Fig. 7). In I4₁₂₂, the distal part of SDD is not visible due to lack of crystal packing at this region and dynamic disorder, not degradation (Supplementary Fig. 8).

To confirm the observed interface in IKK β dimerization, we performed structure-based mutagenesis on residues that bury the most surface area, L654, W655 and L658. These residues are not within the predicted LZ region. Two purified double mutants, L654D/W655D and W655D/L658D, failed to dimerize as shown by the considerable shift in gel filtration positions (Fig. 5c). Furthermore, equilibrium analytical ultracentrifugation experiments showed that WT IKK β is indeed a dimer while the IKK β mutants are either a monomer or have a 167-fold weaker dimerization affinity (Fig. 5c).

Dimerization in IKK β activation but not activity

The geometry of the IKK β dimer with KDs facing away from each other suggests that each IKK β protomer is independent in its kinase activity. To confirm this, we transfected HEK293T cells with hIKK β EE mutants, L654D/W655D (654–5), W655D/L658D (655–8) and L654D/W655D/L658D (654-5-8), and found that all mutants exhibited robust kinase activity. (Fig. 5d). Insect cell expressed and purified dimerization mutants showed the same results. In addition, a purified IKK β construct with truncation into SDD (1–643) that is monomeric in solution (Fig. 5a) was active in phosphorylating I κ B α (Fig. 5d).

Previous studies have shown that IKK β can undergo trans-autophosphorylation and activation upon transfection 13,16. This autophosphorylation and phosphorylation by TAK1 likely both contribute to IKK β activation upon cell stimulation. To determine whether the observed dimerization interface is critical for this activation, we tested dimerization mutations in IKK β activation upon overexpression in HEK293T cells (Fig. 5e). While WT IKK β was robustly activated, the L654D/W655D and W655D/L658D mutants completely and partially lost this activation, respectively, in a manner that correlates with the degree of impairment in dimerization (Fig. 5c, e). Interestingly, overexpression of IKK β in NEMO-deficient mouse embryonic fibroblasts (MEFs) resulted in its activation, but to a lesser extent than in WT MEFs (Fig. 5f). We found that IKK β dimerization mutants failed to interact with NEMO efficiently (Fig. 5g), perhaps due to reduced avidity towards oligomeric NEMO. Therefore, although IKK β kinase activity does not depend on dimerization once its activation loop is phosphorylated, IKK β activation requires dimerization and is likely enhanced by interaction with NEMO.

DISCUSSION

The IKK β structure presented here predicts that IKK α , TBK1 and IKKi all exhibit an overall structural organization that comprises KD, ULD, and SDD. Although an ULD was not predicted in IKK α , conservation of buried residues in this region and of the ULD-SDD interface suggests that IKK α also has this domain (Supplementary Fig. 4). ULD and SDD likely share the common structural role but may have evolved additional, differential functions in the individual kinases. SDD-mediated dimerization may also be conserved. In particular, residues at the observed IKK β dimerization interface are highly conserved in IKK α (Supplementary Fig. 4), explaining that IKK α and IKK β can form both homo- and hetero-dimers 8,13. Given our structure of IKK β and the previously determined structures of NEMO fragments 33–37, we may speculate that the full IKK complex is also a dimer, or a dimer of dimers with a molecular mass of around 270 kD or 540 kD. The apparent 700–900 kD molecular mass of the IKK holoenzyme on gel filtration 5–10 may be due to the elongated nature of NEMO and the complex (Supplementary Fig. 9).

Because the conserved IKK β dimer structure does not place KD close to each other for trans-autophosphorylation, we wondered whether higher order oligomerization, which is likely enhanced by NEMO and its ability to bind ubiquitin 33,34, is responsible for this autoactivation. In both P1 and I4₁22, IKK β exist as dimers of dimers (Fig. 5h, Supplementary Fig. 10). In particular, active sites of two neighboring protomers in the tetramer face each other such that activation loop from one protomer might reach into the active site of the other (Fig. 5i), which may mimic an autophosphorylation state.

The ULD-SDD region of IKK β directly interacts with the C-terminal portion of I κ B α . This interaction may serve several purposes. First, it likely orients I κ B α so that its N-terminal cognate phosphorylation sites are presented to the KD active site (Fig. 4g). Without this interaction, IKK β preferentially phosphorylates the C-terminal PEST motif of I κ B α . Second, the interaction appears to enhance IKK β activity. Third, phosphorylation at the PEST by CK2 or other kinases may regulate I κ B α interaction with IKK β and hence affect phosphorylation at the cognate sites. Fourth, the interaction may allow concerted

phosphorylation at both S32 and S36 of I κ B α without intervening dissociation. In MAP kinase cascades that involve dual phosphorylations, specific docking interactions occur between the kinases and their substrates 27,38,39. The β -catenin protein in the Wnt signaling pathway contains the same dual phosphorylated destruction box motif as I κ B α 40. Consistently, β -catenin is brought to the responsible kinase GSK-3 via the adaptor protein Axin, allowing both specificity and concerted phosphorylation 41,42. Therefore, in a general statement that structure serves the function, the IKK β structure seems to fit perfectly its function as the I κ B α kinase, poised to turn on the important NF- κ B pathway specifically, efficiently and concertedly in response to cellular physiology.

METHODS SUMMARY

Xenopus laevis IKK β was expressed in insect cells and purified to homogeneity using Ni-affinity, ion exchange and gel filtration chromatography. It was crystallized at 4°C in polyethylene glycol (PEG) 6000 and K/Na phosphate for the P1 and I4₁₂₂ crystals, respectively. The structure was determined by multi-wavelength anomalous diffraction (MAD) of the selenomethionyl protein.

Methods

Protein expression and purification

To elucidate the molecular basis of IKK β function, we expressed IKK β from a number of species using baculovirus mediated insect cell expression. The *Xenopus laevis* IKK β (xIKK β) sequence in the NCBI database starts at a Met residue that is equivalent to M17 of both the human and mouse IKK β sequences (hIKK β and mIKK β). Translation of the DNA sequence preceding the ATG codon of M17 revealed sequences that are almost identical to residues 9–16 of hIKK β and mIKK β . These were taken as part of the xIKK β sequence and residues 1–8 were taken from the corresponding mIKK β sequence. This reconstructed xIKK β sequence has the same residue numbering as the hIKK β sequence until after the scaffold/dimerization domain.

Various constructs of IKK β WT and the phosphomimic S177E/S181E mutant were designed with an N-terminal polyhistidine tag and a Tobacco Etch Virus (TEV) protease cutting site between the tag and the protein. Recombinant IKK β baculoviruses were made in DH10BAC cells, amplified, and used to infect Hi5 insect cells in serum-free media (Invitrogen). The cells were cultured in suspension and harvested 48 hrs post-infection. The recombinant proteins were purified by Ni-affinity chromatography, anion-exchange and gel-filtration chromatography. For crystallization, the polyhistidine tag was cleaved by the TEV protease during protein purification.

All I κ B α proteins were expressed in *E. coli* using **pET28a**, **pGEX4T3** and **pET-SUMO** vectors and purified by their respective affinity tags. For kinase assays, the SUMO tag was cleaved off I κ B α proteins. His-ULD and His-ULD-SDD of hIKK β were also expressed in *E. coli* using the **pET28a** vector.

Crystallization and data collection

Unlike many protein kinases, IKK β KD cannot be recombinantly expressed as a well-behaved biochemical entity for structural studies. In addition, after mapping a compact region by limited proteolysis, IKK β was still refractory to crystallization, both alone and in the presence of various ATP analogues. To overcome this obstacle, we used several IKK β inhibitors including Cmpd1 and Cmpd2 (Supplementary Fig. 1), which were identified against the S177E/S181E (EE) mutant, in co-crystallization. A human IKK β (hIKK β) EE construct (residues 1–678) lacking only the C-terminal NBD did crystallize; however, these crystals only diffracted to ~ 7.5 Å resolution. Searching IKK β orthologues that may give better crystals led to a success in crystallizing the analogous region of *Xenopus laevis* IKK β (xIKK β) EE (residues 4–675) (Fig. 1a).

The xIKK β (S177E/S181E) protein containing residues 4–675 was concentrated by ultrafiltration (Amicon) to about 15 mg/ml in 20 mM Tris-HCl (pH 8.0), 150 mM NaCl and 10 mM dithiothreitol (DTT). It was mixed with an inhibitor compound in 1:2 molar ratios before crystallization. Cmpd1 is (4-((4-(4-(chlorophenyl)pyrimidin-2-yl)amino)phenyl)(4-(2-hydroxyethyl)piperazin-1-yl)methanone and Cmpd2 is 1-(4-(4-((4-(4-(pyridin-4-ylsulfonyl)phenyl)pyrimidin-2-yl)amino)benzoyl)piperazin-1-yl)ethanone. The P1 crystals were grown using hanging drop vapor diffusion at 4°C by mixing equal volumes of the purified protein and the crystallization condition of 100 mM N-(2-acetamido) iminodiacetic acid at pH6.5, 10 % (w/v) polyethylene glycol (PEG) 6000, 50 mM Li₂SO₄, 300 mM NaCl and 10 mM DTT. The I4₁22 crystals were grown at 4°C with well solution containing 1.8 M K/Na phosphate at pH 5.6 and 10 mM DTT. For data collection, all crystals were flash frozen in the respective crystallization conditions supplemented with 25 % (v/v) ethylene glycol. Diffraction data were collected at the 24ID-C beam line of the Advanced Photon Source (APS). Multi-wavelength anomalous diffraction (MAD) data of heavy-atom derivative crystals or selenomethionyl (Se-Met) crystals were collected near the respective absorption edges. All diffraction data were processed using the HKL2000 suite 43 and their statistics are shown in Supplementary Table 1 and Supplementary Table 2.

Structure determination, refinement and analysis

The initial xIKK β crystals grew in the P1 space group in the presence of the inhibitor Cmpd1 or Cmpd2 and diffracted to 3.6 Å resolution. Selenomethionyl crystals were obtained, but we failed to locate the large number of expected Se sites. Among the extensive heavy atom searches, a Yb-derivative was obtained, with 8 well-defined sites, which likely correspond to 8 IKK β molecules in the asymmetric unit. However, the electron density map calculated from a 3-wavelength Yb-anomalous diffraction data set was insufficient for tracing and phase combination with the selenomethionyl data set could not be performed due to non-isomorphism.

The structure determination was eventually successful in the alternative crystal form of I4₁22, which contains 1 molecule of IKK β in complex with Cmpd1 and diffracted to 4.0 Å resolution, using multi-wavelength anomalous diffraction (MAD) of the selenomethionyl crystals (Supplementary Table 1 and 2, Supplementary Fig. 2). Twelve Se-sites were determined using the program Shelx D 44 and refined with the program MLPHARE in the

CCP4 suite 45. MAD phases were calculated at 4.0 Å resolution with data from I4₁22 crystals using the program SHARP 46. A few cycles of model building and refinement were carried out with the program O 47 and REFMAC with TLS parameterization 48. The I4₁22 crystals contain one monomer in the asymmetric unit and 80% solvent when calculated with the entire IKK β construct and 84% solvent when considering only the ordered part of the structure. The inhibitor Cmpd1 has density in the MAD experimental map and the Fo-Fc omit map. The Dundee PRODRG2 Server was used to generate topology and restraint files of the compound for refinement. The refined model contains residues 16–236, 243–286, 290–376, 384–394, 401–475 and 528–637 and Cmpd1.

The structure of the P1 form was determined by molecular replacement using the refined model of the I4₁22 crystal form as the search model, in which 8 molecules were located. Sites of the SAD data set of a selenomethionyl crystal in the P1 space group was calculated by difference Fourier and used to verify residue registration in the P1 structure. Refinement in the P1 structure was conducted at 3.6 Å resolution and incorporated tight non-crystallographic symmetry (NCS) restraints (0.02 Å rmsd in atom positions). Upon several rounds of refinement at 3.6 Å resolution, new electron densities appeared in the P1 crystal form to complete the model building. Although Cmpd2 was in the crystallization condition, it did not have clear density and was not included in the refinement. The refined model of the P1 crystal form contains four IKK β dimers in the asymmetric unit. Three of the dimers encompass residues 16–236, 243–286, 290–376, 384–394, 401–551 and 559–666. One dimer contains the same residues as the structure in I4₁22. The structures were analyzed using the CCP4 suite 45 and the Dali server 49, and the figures were made using PyMOL 50.

GST-pulldown

The tagged proteins were first purified with glutathione or Ni-NTA beads and their expression levels were assessed by SDS-PAGE. Beads containing estimated equivalent quantities of the tagged proteins were mixed with the cell lysates or the purified versions of the interaction partners. The mixtures were incubated at room temperature for 1 hr with rotation. After centrifugation, the supernatants were removed. The beads were then washed twice, eluted and subjected to SDS-PAGE analysis. All pulldown experiments were repeated two to four times with consistency.

Kinase assays using anti-phospho-I κ B α antibody

The hIKK β proteins (0.1 μ g/ μ l) were incubated with recombinant I κ B α (1 μ g/ μ l) in 50 mM Tris-HCl at pH 8.0, 100 mM NaCl, 10 mM MgCl₂, and 2 mM DTT for 30 min at 30°C. SDS-PAGE sample buffer was used to terminate the reactions. The products were separated on 15 % SDS-PAGE and transferred to PVDF membranes. Anti-phospho-I κ B α antibody (Cell Signaling Technology, Inc.) was used to detect phospho-I κ B α .

Determination of K_m of hIKK β for I κ B α (1–54) and full-length

To derive the K_m of hIKK β for full-length I κ B α , kinase assays were performed at substrate concentrations of 2.8 μ M, 5.3 μ M, 10.6 μ M, 21.3 μ M, 42.5 μ M, and 85 μ M. Similarly, to derive the K_m of hIKK β for I κ B α (1–54), kinase assays were performed at substrate

concentrations of 5.3 μM , 10.6 μM , 21.3 μM , 42.5 μM , 85 μM , and 170 μM . A time course of the kinase reactions was first performed to select a hIKK β amount and a time point within which the reactions are linear with time. The final selected reactions contain 10 ng baculovirus-expressed hIKK β , 100 mM cold ATP and 1 μl [γ - ^{32}P]ATP (3000Ci/mmol, 1mCi/100 μl) in 25 μl reaction buffer containing 50 mM Tris-HCl at pH 8.0, 100 mM NaCl, 10 mM MgCl_2 and 2 mM DTT. The phospho-transfer reaction was allowed to proceed for 10 minutes at 30 $^\circ\text{C}$ and quenched with SDS-PAGE sample buffer. The products were separated on 15 % SDS-PAGE and subjected to autoradiography. The relative amounts of phosphorylated I κ B α were quantified using ImageJ, plotted against total I κ B α concentrations and fitted using non-linear regression to the Michaelis-Menten equation to obtain the K_m using SigmaPlot.

Transfection, immunoprecipitation and kinase assay

The constructs Flag-hIKK β EE and its truncation mutants, HA-hIKK β EE and its dimerization mutants L654D/W655D, W655D/L658D and L654D/W655D/L658D, Flag-hIKK β and its dimerization mutants L654D/W655D and W655D/L658D were generated in the vector **pcDNA3** by conventional PCR. All IKK β constructs were transfected in HEK293T cells with Lipofectamine 2000 (Invitrogen). After 24 hrs, cell extracts were immunoprecipitated with anti-Flag antibodies bound to agarose beads (M2, Sigma) or anti-HA bound to agarose beads (Sigma). IKK β kinase assays were essentially done as described 6,13. Briefly, immunoprecipitates were incubated with 2 μM full-length I κ B α (1–317) in 20 mM HEPES at pH 7.5, 10 mM MgCl_2 , 20 mM β -glycerophosphate, 10 mM PNPP, 50 mM Na_3VO_4 , 1 mM DTT, 20 mM ATP, and (1–10 mCi) [γ - ^{32}P]ATP at 30 $^\circ\text{C}$ for 30 min, and subjected to SDS-PAGE and autoradiography. Immunoblotting was performed using anti-Flag (Sigma), anti-HA (Sigma) or anti-IKK β antibodies (Upstate, 05–535).

Equilibrium analytical ultracentrifugation measurements

Experiments were performed in a Beckman XL-A/I analytical ultracentrifuge (Beckman-Coulter, Palo Alto CA, USA), utilizing six-cell centerpieces with straight walls, 12 mm path length and sapphire windows. Samples were kept and diluted in 50 mM Tris-HCl at pH 8.0 and 300 mM NaCl. Samples from wild-type protein were diluted to 6.9, 4.5 and 2.4 μM , mutant L654D/W655D was diluted to 7.4, 4.8, and 2.6 μM and mutant W655D/L658D to 4.9, 3.2, and 1.7 μM for channels A, B and C, respectively. Dilution buffer was used as blank. All samples were run at 4 $^\circ\text{C}$ at 9000 rpm (held for 20 h, then four scans with 1 h interval), 11000 rpm (held 10 h then four scans with 1 h interval), 14000 rpm (held 10 h then four scans with 1 h interval), 17000 rpm (held 10 h then four scans with 1 h interval). Detection was by UV at 280 nm. Solvent density and protein v -bar at both temperatures were determined using the program SednTerp (Alliance Protein Laboratories, Corte Cancion, Thousand Oaks, CA, USA). For calculation of K_D and apparent molecular weight, all useful data were used in a global fit, using the program HeteroAnalysis, obtained from University of Connecticut (www.biotech.uconn.edu/auf).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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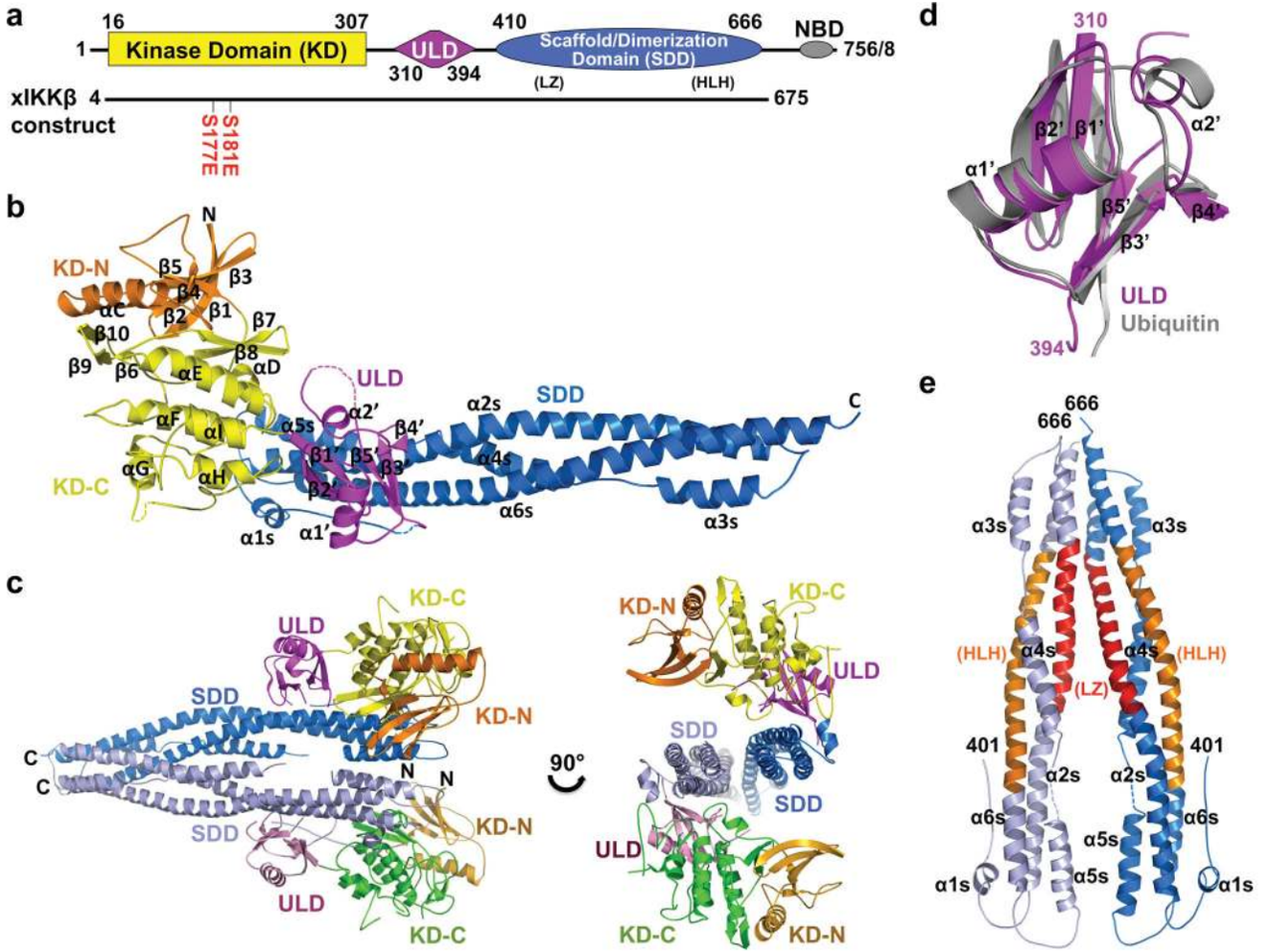


Figure 1. Structure of xIKKβ

a, Linear representation of IKKβ showing the boundaries for the kinase domain (KD), the ubiquitin-like domain (ULD) and the scaffold/dimerization domain (SDD). Sequences of hIKKβ and xIKKβ are of 756 and 758 residues, respectively, differing only at the most C-terminal region. The crystallized xIKKβ construct is shown. The previously designated leucine zipper (LZ) and helix-loop-helix (HLH) regions are shown in parentheses. **b**, Ribbon diagram of an xIKKβ protomer in the P1 crystal form. The N- and C-termini, KD N-lobe (orange) and C-lobe (yellow), ULD (magenta) and SDD (blue) are labeled. Secondary structural elements are labeled, with those in ULD followed by (') and those in SDD followed by (s). **c**, Ribbon diagram of an xIKKβ dimer. **d**, Superposition of ULD (magenta) with ubiquitin (gray). **e**, Ribbon diagram of an SDD dimer, showing locations of the previously designated LZ (red) and HLH (orange) regions.

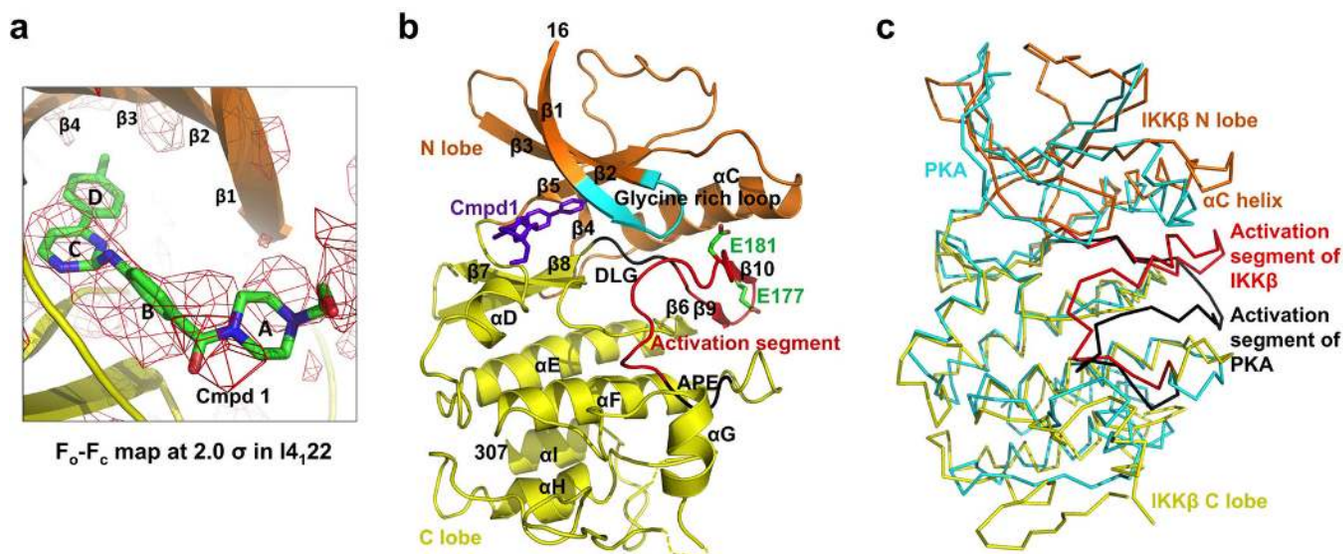


Figure 2. Inhibitor bound xIKK β kinase domain (KD)

a, F_o-F_c electron density map for Cmpd1 in the I4₁₂₂ structure, contoured at 2.0σ . Carbon, nitrogen and oxygen atoms are shown in green, blue and red, respectively. **b**, Structure of xIKK β KD. Glycine-rich loop: cyan; activation segment: red except that the DLG and APE motifs are in black; Cmpd1: purple. Side chains of phosphomimic residues E177 and E181 are shown. **c**, Superposition between xIKK β (orange and yellow) and PKA (cyan, PDB code 1ATP). The activation segments of xIKK β and PKA are shown in red and black, respectively.

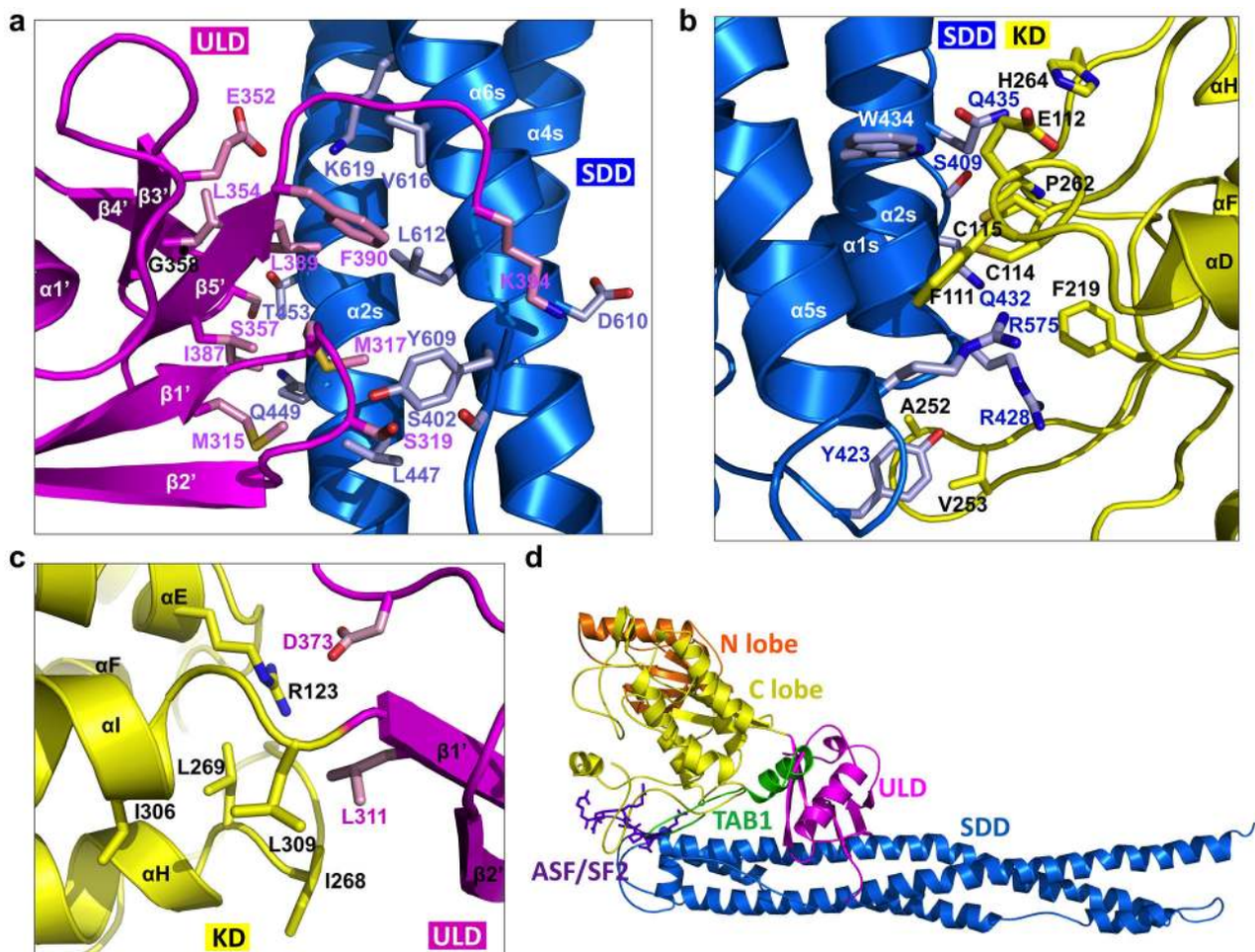


Figure 3. Interactions among the KD, the ULD and the SDD

a, Interaction between ULD (magenta) and SDD (blue). Important interfacial side chains are shown with nitrogen atoms in blue, oxygen atoms in red, sulfur atoms in orange, and carbon atoms in either pink (for ULD) or light blue (for SDD). Location of G358 is marked with a black ball on the main chain. **b**, Interaction between KD (yellow) and SDD (blue). **c**, Interaction between KD (yellow) and ULD (magenta). **d**, Locations of the TAB1 peptide (green ribbon, PDB code 2EVA) and the ASF/SF2 peptide (purple stick model, PDB code 1WBP) relative to the IKK β structure after superposition of the corresponding kinase domains.

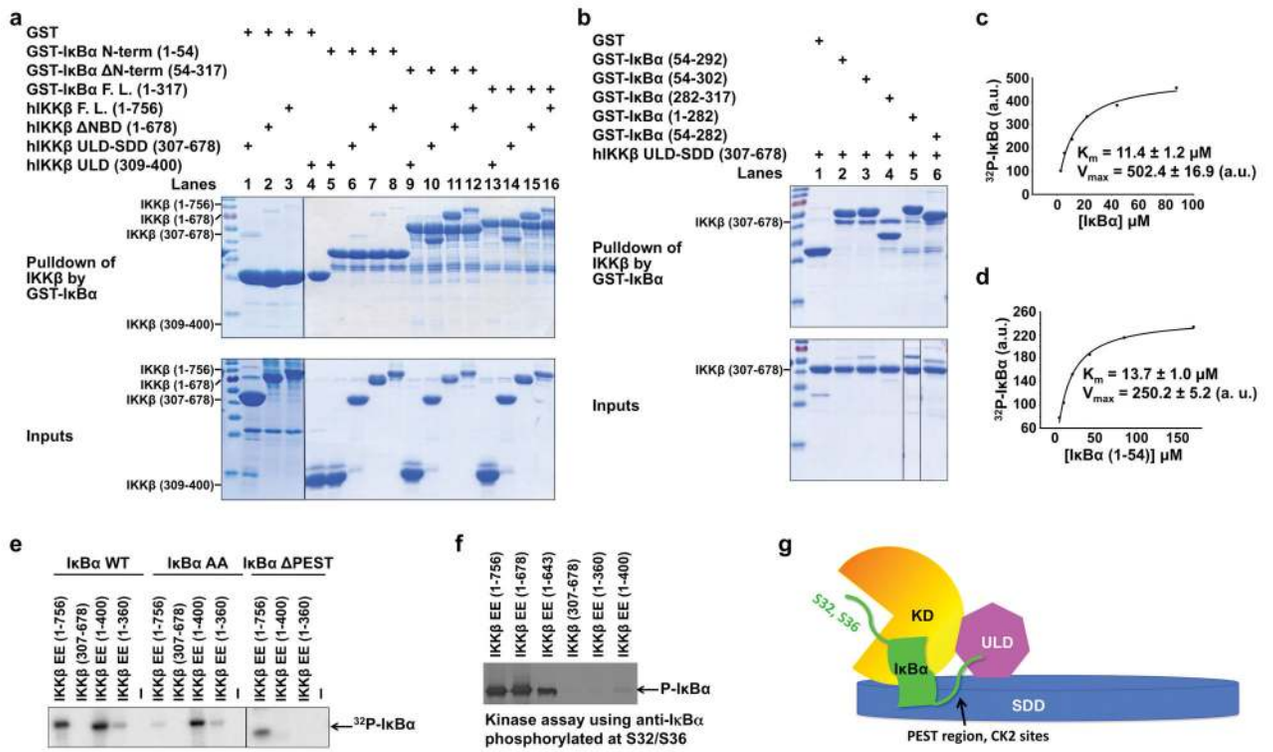


Figure 4. ULD-SDD restricts IKKβ specificity while ULD is required for catalytic activity
a, b, Pull-down of hIKKβ constructs using GST-IκBα constructs, showing the reciprocal interaction between ULD-SDD of IKKβ and C-terminal region of IκBα containing ankyrin repeats and PEST region. **c, d**, Measurement of K_m and relative V_{max} of IKKβ against full-length IκBα (c) and the N-terminal region of IκBα (1–54) (d). a.u.: arbitrary unit. **e**, Kinase assay of purified hIKKβ proteins against IκBα, its S32A/S36A mutant (AA), or its PEST-deletion construct (ΔPEST, 1–282) using $[\gamma\text{-}^{32}\text{P}]\text{ATP}$. **f**, Kinase assay of purified hIKKβ proteins using antibody against IκBα phosphorylated at S32 and S36. **g**, A schematic model showing that the interaction between SDD of IKKβ and C-terminal region of IκBα may position the N-terminal cognate phosphorylation sites of IκBα to the active site of IKKβ.

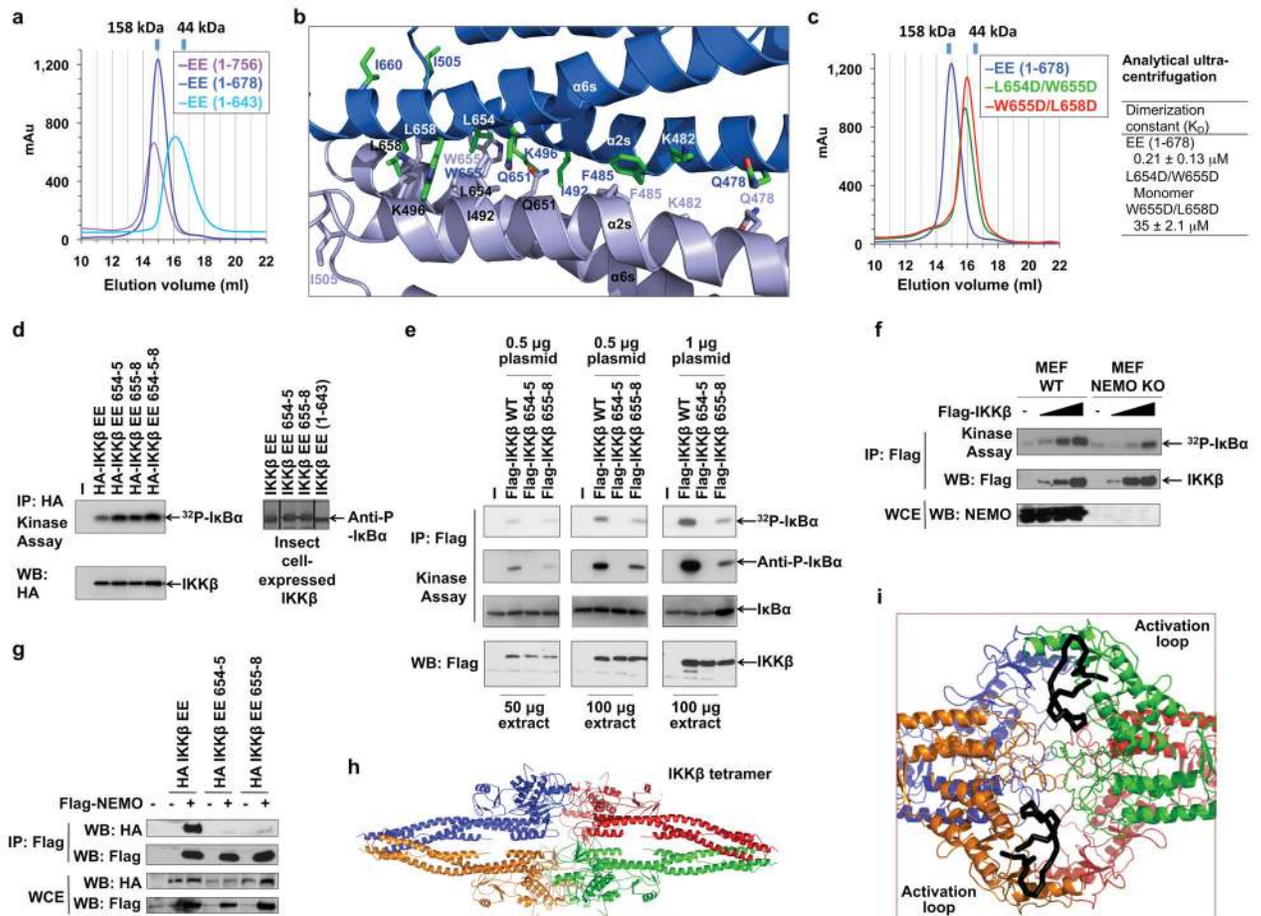


Figure 5. Dimerization is critical for IKK β activation but not for its activity

a, Gel filtration profiles of various hIKK β constructs showing that those containing the KD-ULD-SDD region (1–756, 1–678) are dimeric while a further truncated construct (1–643) is monomeric. **b**, The dimerization interface of α IKK β . **c**, Structure-based mutations disrupt hIKK β dimerization as shown by gel filtration and analytical ultracentrifugation. **d**, Kinase activity of HEK293T cell transfected or insect cell purified hIKK β EE and dimerization defective mutants L654D/W655D (654–5), W655D/L658D (655–8) and L654D/W655D/L658D (654-5-8). **e**, Autoactivation of HEK293T cell transfected hIKK β WT and its dimerization defective mutants. **f**, Transfection of hIKK β into WT and NEMO $^{-/-}$ MEFs, showing reduced IKK β activation in the absence of NEMO. **g**, Dimerization mutants of IKK β showed reduced interaction with NEMO. **h**, A tetramer of α IKK β in the P1 structure. **i**, A close-up view of the tetramer interface, showing that the activation loops of neighboring protomers (black) face each other.