The nuclear protein IkBζ Forms a Transcriptionally Active Complex with Nuclear Factor-κB (NF-κB) p50 and the Lcn2 Promoter via the N- and C-terminal Ankyrin Repeat Motifs

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The nuclear protein IkBζ, comprising the N-terminal trans-activation domain and the C-terminal ankyrin repeat (ANK) domain composed of seven ANK motifs, activates transcription of a subset of nuclear factor-κB (NF-κB)-dependent innate immune genes such as Lcn2 encoding the antibacterial protein lipocalin-2. Lcn2 activation requires formation of a complex containing IkBζ and NF-κB p50, a transcription factor that harbors the DNA-binding Rel homology region but lacks a trans-activation domain, on the promoter with the canonical NF-κB-binding site (κB site) and its downstream cytokine-rich element. Here we show that IkBζ productively interacts with p50 via Asp-451 in the N terminus of ANK1, a residue that is evolutionarily conserved among IkBζ and the related nuclear IkB proteins Bcl-3 and IkBNS. Threonine substitution for Asp-451 abrogates direct association with the κB-site-binding protein p50, complex formation with the Lcn2 promoter DNA, and activation of Lcn2 transcription. The basic residues Lys-717 and Lys-719 in the C-terminal region of ANK7 contribute to IkBζ binding to the Lcn2 promoter, probably via interaction with the cytokine-rich element required for Lcn2 activation; glutamate substitution for both lysines results in a loss of transcriptionally active complex formation without affecting direct contact of IkBζ with p50. Both termini of the ANK domain in Bcl-3 and IkBNS function in a manner similar to that of IkBζ to interact with promoter DNA, indicating a common mechanism in which the nuclear IkBs form a regulatory complex with NF-κB and promoter DNA via the invariant aspartate in ANK1 and the conserved basic residues in ANK7.

Nuclear factor-κB (NF-κB) plays central roles in host defense and inflammation as a homo- or heterodimer of NF-κB/Rel family proteins by controlling the expression of genes for pro-inflammatory cytokines, chemokines, and anti-bacterial proteins (1–4). The mammalian NF-κB family is composed of five structurally related polypeptides: p50, p52, p105 (the precursor of p50), p100 (the precursor of p52), p65 (also known as RelA), RelB, and c-Rel. They share the Rel homology region, which mediates dimerization, nuclear translocation, binding to specific DNA sequences known as NF-κB-binding elements (κB sites), and association with one of the IkB family proteins (1–4). Among the members of the family, p65, RelB, and c-Rel have an ability to activate transcription by themselves via the C-terminal trans-activation domain, which is absent in the smaller p50 and p52 proteins. In resting cells, NF-κB dimers are retained in the cytoplasm by associating with a member of the prototypical/cytoplasmic IkB proteins including IkBa, IkBβ, and IkBe (1–4). Cell activation with appropriate stimuli such as bacterial LPS leads to phosphorylation-induced degradation of cytoplasmic IkBs and resultant liberation of NF-κB dimers (1, 5, 6). The released NF-κB dimers subsequently translocate to the nucleus and thus induce the expression of primary response genes via binding to κB sites on their promoter/enhancer regions (1–4).

The primarily induced gene products include the atypical/nuclear IkB proteins Bcl-3 (7, 8), IkBζ (also known as MAIL or INAP) (9–11), and IkBNS (12), which are not expressed in resting cells. In contrast to the cytoplasmic IkBs, which preferentially associate with NF-κB dimers that possess at least one p65 or c-Rel subunit such as the p50-p65 heterodimer (13), the nuclear IkBs prefer the p50 (or p52) homodimer as a partner to directly regulate the activation of secondary response genes for appropriate host defense responses (14–19). Thus, a part of the secondary response genes is controlled by NF-κB-dependent induction of nuclear IkBs and their subsequent association with NF-κB. For instance, the secondarily induced genes Lcn2 (encoding the antibacterial protein lipocalin-2) and Ptx3 (encoding the antibacterial protein pentraxin 3) are activated by the nuclear IkB protein IkBζ (20–24), which serves as a key regulator in the immune system (16, 18, 25, 26). On the other hand, Bcl-3 participates in transcriptional control of the chemokine-encoding genes Ccl2 for monococyte chemoattractant protein-1 (MCP-1) and Cxcl10 for interferon-γ-induced protein 10 (IP-10) (27), and IkBNS regulates transcription of Il6 (encoding the pro-inflammatory cytokine interleukin-6) and Il12b (encoding interleukin-12 subunit p40) (28, 29).

The IkB proteins are characterized by the presence of the ankyrin repeat domain (ARD). The ARD of the cytoplasmic or nuclear IkBs contains six or seven ankyrin repeat (ANK) motifs, respectively. The ANK is an evolutionarily conserved protein...
motif of about 33 amino acid residues that forms an L-shaped structure comprising a β hairpin and two antiparallel α helices. Consecutive ANK motifs generally stack together to serve as an underlying architecture of a modular specific protein-interacting interface (30–32). The ARD of IκBα mediates the association with NF-κB dimers via direct binding to the NF-κB Rel homology region (23, 33, 34). Although the ARD of nuclear IκBα displays a high sequence similarity to that of cytoplasmic IκBs except for the additional seventh motif ANK7 (10, 12, 35, 36), the preference for NF-κB dimer species differs between the nuclear and cytoplasmic IκBs, as described above (7–9, 12, 16, 37). Furthermore, there exists a difference in interaction with κB DNA. Although IκBα promotes the dissociation of a p65-containing dimer from the promoter DNA (38), the nuclear IκBs are generally assumed to interact indirectly with the κB site via binding to NF-κB p50 or p52 (7, 8, 23, 33, 34). In addition, we have recently shown that IκBζ interaction with the Lcn2 promoter also requires a region downstream of the κB site (23). However, the molecular mechanism underlying assembly of the nuclear IκB-containing regulatory complex has not been well elucidated.

In the present study, we show that IκBζ, containing the N-terminal trans-activation domain and the C-terminal ARD, forms a transcriptionally active complex with the Lcn2 promoter via both Asp-451-mediated association with the κB-site-binding protein p50/p52 and Lys-717/Lys-719-dependent interaction with the downstream extra κB site on the Lcn2 promoter. Asp-451, an invariant residue among the nuclear IκBs, is present in the N terminus of ANK1; Lys-717 and Lys-719 exist in the region C-terminal to the second α helix of ANK7, and the corresponding sites are also occupied by basic residues in Bcl-3 and IκBNS. Both termini of the ARD in these proteins serve in a manner similar to that of IκBζ, indicative of a common mechanism by which nuclear IκBs form a p50/p52-containing regulatory complex on target gene promoters.

Results

The Invariant Aspartate in ANK1 of IκBζ and Other Nuclear IκBs Is Crucial for Interaction with NF-κB—It is known that the ARD of the nuclear IκB proteins IκBζ, IκBNS, and Bcl-3 (Fig. 1A) is responsible for binding to NF-κB p50 (33, 34). To determine the N-terminal boundary of the IκBζ region required for interaction with p50, we expressed and purified a series of N-terminally truncated IκBζ as GST-fused proteins and tested their ability to bind to p50. As shown in Fig. 1B, IκBζ-(449–728) fully interacted with p50, as did IκBζ-(414–728). On the other hand, IκBζ-(453–728) failed to bind to p50 (Fig. 1B), suggesting a role for amino acid residues 449–452. Among the four residues, Asp-451 in ANK1 is the only one that is completely conserved during evolution (Fig. 1A). Intriguingly, the aspartate also exists in the other nuclear IκB proteins IκBNS and Bcl-3 (Asp-60 and Asp-127, respectively) but is replaced by threonine or serine in cytoplasmic IκB members (Fig. 1A). Consistent with the conservation, the replacement of Asp-451 by threonine impaired IκBζ binding to p50 in a GST pulldown assay (Fig. 1C). The D451T substitution in IκBζ also resulted in a loss of its co-immunoprecipitation with p50 when FLAG-tagged full-length IκBζ and HA-p50 were expressed in HEK293T cells (Fig. 1D). Thus, the invariant residue Asp-451 in ANK1 plays a crucial role in IκBζ interaction with p50.

We next investigated the role of the corresponding aspartate residue in other nuclear IκB proteins (Asp-60 in IκBNS and Asp-127 in Bcl-3). Threonine substitution for Asp-60 in IκBNS led to an impaired interaction with p50 both in a GST pulldown assay using purified proteins (Fig. 1E) and in a co-immunoprecipitation assay using proteins expressed in HEK293T cells (Fig. 1F). Similarly, compared with wild-type Bcl-3, a mutant protein with threonine substitution for Asp-127 interacted with p50 much less efficiently both in vitro (Fig. 1B) and in vivo (Fig. 1G). These findings highlight a conservative role for the invariant aspartate of ANK1 in direct interaction of nuclear IκBs with p50.

Nuclear IκB proteins are also known to form a complex with NF-κB p52, a protein homologous to p50 (7, 12, 37). As expected from the homology, in vitro complex formation with p52 was impaired by threonine substitution for the invariant aspartate in ANK1 of nuclear IκBs: Asp-451 in IκBζ, Asp-60 in IκBNS, and Asp-127 in Bcl-3 (Fig. 2A). The critical role for the aspartates was confirmed by co-precipitation of p52 with IκBζ (Fig. 2B), IκBNS (Fig. 2C), and Bcl-3 (Fig. 2D) when ectopically expressed in HEK293T cells. Thus, p52 likely interacts with nuclear IκB proteins in a manner similar to the way p50 does.

Asp-451 of IκBζ Is Involved in Transcriptional Activation—As we have shown previously (23), expression of IκBζ along with p50 results in transcriptional activation of the promoter of the lipocalin-2-encoding gene Lcn2 in p50/-/IκBζ-deficient mouse embryonic fibroblasts (MEFs) (Nfkbi−/−; Nfkbez−/− MEFs) (Fig. 3A). To examine the role for Asp-451 of IκBζ in p50-dependent Lcn2 activation, we used p50/-/IκBζ-deficient MEFs in which a luciferase reporter is regulated by the Lcn2 promoter (23). As shown in Fig. 3A, IκBζ (D451T), defective in association with p50, activated the Lcn2 promoter much less effectively than the wild-type protein even in the presence of p50. We next tested the role for Asp-451 of ANK1 in IκBζ-mediated activation of the endogenous Lcn2 gene. As shown in Fig. 3B, exogenous expression of wild-type IκBζ in IκBζ-deficient bone marrow-derived macrophages (BMMs) resulted in time-dependent activation of the endogenous Lcn2 gene in response to LPS. On the other hand, IκBζ (D451T), a mutant protein defective in direct interaction with NF-κB p50 (Fig. 1), failed to activate the endogenous Lcn2 gene, although IκBζ (D451T) was expressed at a level similar to that of the wild-type protein (Fig. 3B). Thus, the interaction between IκBζ and p50 appears to be involved in transcriptional activation of Lcn2.

We also investigated the function of Bcl-3 Asp-127, a residue that corresponds to Asp-451 in IκBζ and is crucial for binding to p50 and p52 (Figs. 1 and 2), in gene activation. As shown in Fig. 3C, Bcl-3 (D127T) activated a reporter of the Ccl2 promoter in LPS-stimulated RAW264.7 cells that ectopically expressed p52 but to a significantly lesser extent than the wild-type protein. Thus, Bcl-3 binding to p52 plays a role in Ccl2 activation. On the other hand, IκBζ did not activate Lcn2 in combination with p52 under conditions where p50 fully supported IκBζ (Fig. 3D). This raises the possibility that p52 is incapable of directly interacting with the Lcn2 promoter DNA, although p52 binds to IκBζ as p50 does (Fig. 2B). To address this...
question, we analyzed the formation of the protein-promoter complex using a DNA-binding assay in which DNA bound to a tagged protein is pulled down with tag affinity beads and amplified by PCR (for details, see “Experimental Procedures”). As shown in Fig. 3E, the Lcn2 promoter efficiently interacted with wild-type p50 but not with p50 (Y57A/E60D), a mutant protein defective in binding to the xB site (23), or p56. In contrast to p50, wild-type p52 failed to directly bind to the Lcn2 promoter (Fig. 3E). Thus, 1kBζ appears to activate Lcn2 by specifically interacting with p50.

Lys-717 and Lys-719 in ANK7 of 1kBζ Participate in Lcn2 Activation—We next studied the role for the C-terminal region of the 1kBζ ARD, which comprises seven ANK repeats (Fig. 4A).

For this purpose, we expressed a series of C-terminally truncated 1kBζ proteins in Nfkbi−/−•Nfbiz−/− MEFs to test their ability to activate the Lcn2 promoter. As shown in Fig. 4B, 1kBζ(1–721) was as active as full-length 1kBζ of 728 amino acids. By contrast, Lcn2 was activated much more weakly by 1kBζ(1–716) (Fig. 4C) than by 1kBζ(1–718) and only marginally by 1kBζ(1–716) (Fig. 4B), indicating a crucial role for the C-terminal region of 1kBζ ANK7 (amino acids 717–721) in vivo. GST-fused 1kBζ (C), 1kBζ(E), or Bcl-3 (E) with or without the indicated amino acid substitution was incubated with His-p50, followed by analysis as in B, D, and F, and the role of 1kBζ Asp-451, 1kBζ Asp-60, and Bcl-3 Asp-127 in interaction with NF-κB p50 in vivo. FLAG-tagged 1kBζ (D), 1kBζ (F), or Bcl-3 (G) with or without the indicated amino acid substitution was co-expressed with HA-p50 in HEK293T cells, and proteins in the cell lysate were immunoprecipitated (IP) with anti-FLAG antibody, followed by immunoblot analysis with the indicated antibody (Blot). Positions for marker proteins are indicated in kilodaltons.

FIGURE 1. Asp-451 in ANK1 of 1kBζ and the corresponding residues of 1kBζ, Bcl-3 and B-3 are involved in interaction with NF-κB p50. A, domain organization of mouse 1kBζ and comparison of the amino acid sequences of 1kBζ and its N-terminally flanking region. Mouse 1kBζ of 728 amino acids contains the nuclear localization signal (NLS), the trans-activation domain (TAD), and the ARD comprised of seven ANK motifs. The amino acid sequences of ANK1 and its N-terminally flanking region of 1kBζ from various species and those of other mouse 1kB proteins are aligned. Mm, Mus musculus; Hs, Homo sapiens; Gg, Gallus gallus; Xt, Xenopus tropicalis; Dr, Danio rerio. B, the 1kBζ N-terminal boundary required for interaction with NF-κB p50. GST-fused 1kBζ with the indicated truncation was incubated with His-p50 and pull-down with glutathione-Sepharose-4B beads, followed by SDS-PAGE analysis with CBB staining. MW, molecular weight. C and E, the role for 1kBζ Asp-451, 1kBζ Asp-60, and Bcl-3 Asp-127 in interaction with NF-κB p50 in vitro. GST-fused 1kBζ (C), 1kBζ (E), or Bcl-3 (E) with or without the indicated amino acid substitution was incubated with His-p50, followed by analysis as in B, D, and F, and the role of 1kBζ Asp-451, 1kBζ Asp-60, and Bcl-3 Asp-127 in interaction with NF-κB p50 in vivo. FLAG-tagged 1kBζ (D), 1kBζ (F), or Bcl-3 (G) with or without the indicated amino acid substitution was co-expressed with HA-p50 in HEK293T cells, and proteins in the cell lysate were immunoprecipitated (IP) with anti-FLAG antibody, followed by immunoblot analysis with the indicated antibody (Blot). Positions for marker proteins are indicated in kilodaltons.
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**FIGURE 2.** Asp-451 in ANK1 of IkBζ and the corresponding residue of IkBns and Bcl-3 are involved in interaction with NF-κB p50. A, the role for IkBζ Asp-451, IkBns Asp-60, and Bcl-3 Asp-127 in interaction with NF-κB p50 in vitro. GST-fused IkBζ, IkBns, and Bcl-3 with or without the indicated amino acid substitution were incubated with His-p52(1–341), and pulled down with glutathione-Sepharose-4B beads, followed by SDS-PAGE analysis with CBB staining. MW, molecular weight. B–D, the role of IkBζ Asp-451, IkBns Asp-60, and Bcl-3 Asp-127 in interaction with NF-κB p50 in vivo. FLAG-tagged IkBζ (B), IkBns (C), or Bcl-3 (D) with or without the indicated amino acid substitution was co-expressed with HA-p52 in HEK293T cells, and proteins in the cell lysate were immunoprecipitated (IP) with the anti-FLAG antibody, followed by immunoblot analysis with the indicated antibody (Blot). Positions for marker proteins are indicated in kilodaltons.

contains the basic residues Lys-717 and Lys-719, both of which are evolutionarily well conserved (Fig. 4A). The Lcn2 promoter was activated by a mutant IkBζ carrying substitution of the acidic residue glutamate for Lys-717, but to a lesser extent than the wild-type protein (Fig. 4D), and double glutamate substitution for Lys-717 and Lys-719 led to an almost complete loss of Lcn2 activation (Fig. 4D) without affecting the ability to directly interact with p50 (Fig. 4E). Furthermore, LPS-induced activation of the endogenous Lcn2 gene was not observed in BMMs expressing a mutant IkBζ with the K717E/K719E substitution (Fig. 3B). In contrast, simultaneous replacement of Lys-717 and Lys-719 by the other basic residue arginine hardly affected Lcn2 activation (Fig. 4F). These observations imply a possible role for the positive charge at amino acid positions 717 and 719 in IkBζ ANK7.

Lys-717 and Lys-719 in ANK7 of IkBζ Participate in Complex Formation with the Lcn2 Promoter and Not via p50—It should be noted that the basic residues lysine and arginine are both capable of not only forming a salt bridge with a phosphate group of DNA but also of directly interacting with a base of DNA, especially guanine (39, 40). Furthermore, we have shown recently that a cytosine-rich region downstream of the κB site is involved in formation of the active three-species complex IkBζ-p50-DNA (23), suggesting that the extra-κB site makes a direct contact with p50, IkBζ, or both. It seems thus possible that the arginine-replaceable residues Lys-717 and Lys-719 in ANK7 participate in IkBζ interaction with the Lcn2 promoter DNA. To test this possibility, we analyzed the in vitro formation of the IkBζ-p50-DNA complex using the method used in Fig. 3E. Under conditions where the Lcn2 promoter fragment was fully co-precipitated with wild-type IkBζ, the co-precipitation was not caused by IkBζ (D451T), defective in direct binding to p50 (Fig. 5A). The D451T substitution also disrupted IkBζ interaction with the endogenous promoter of Lcn2, as shown by ChIP analysis using cells that stably expressed the wild-type or mutant protein (Fig. 5B). Further analysis with the anti-RNA polymerase II antibody revealed that the impairment of IkBζ-p50 association resulted in loss of the in vivo formation of a transcriptionally active complex on the Lcn2 promoter (Fig. 5B). The requirement of IkBζ-p50 association for interaction with the promoter is in good agreement with the following observation. Both p50 protein and the p50-binding site on the promoter (the κB site) were required for IkBζ interaction with DNA (Fig. 5A). Thus, Asp-451 indirectly participates in complex formation with the Lcn2 promoter via direct binding to p50. On the other hand, Lys-717 and Lys-719 in IkBζ may directly interact with target DNA because the K717E/K719E substitution abolished not only the recruitment to the Lcn2 endogenous promoter for subsequent active complex formation (Fig. 5B) but also the in vitro interaction with the target DNA without affecting binding to p50 (Fig. 5A). As indicated from the finding that arginine can replace Lys-717 and Lys-719 in Lcn2 activation (Fig. 4F), the replacement did not affect active complex formation with the Lcn2 promoter (Fig. 5D). These findings indicate that IkBζ interacts with the Lcn2 promoter via the arginine-replaceable residues Lys-717 and Lys-719 in a manner independent of direct binding to p50.
Furthermore, proline substitution for the evolutionarily well conserved residue Gly-718, which locates between Lys-717 and Lys-719 in IκBζ ANK7 (Fig. 4A), resulted in a loss of both interaction with the Lcn2 promoter (Fig. 5A) and Lcn2 activation (Fig. 5C). The observation suggests that correct orientation of Lys-717 and Lys-719 toward the target DNA may be required for association of IκBζ to the Lcn2 promoter. On the other hand, neither S720A nor I721S substitution affected Lcn2 activation (Fig. 5C).

Acetylation of IκBζ Does Not Seem to Be Involved in Lcn2 Activation—The significance of Lys-717 and Lys-719 of IκBζ toward the target DNA may be required (Fig. 5). The conservation of Lys-717 and Lys-719 in IκBζ ANK7 (Fig. 4A), resulted in a loss of both interaction with the Lcn2 promoter (Fig. 5A) and Lcn2 activation (Fig. 5C). The observation suggests that correct orientation of Lys-717 and Lys-719 toward the target DNA may be required for association of IκBζ to the Lcn2 promoter. On the other hand, neither S720A nor I721S substitution affected Lcn2 activation (Fig. 5C).

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These observations suggest that acetylation of Lys-717 or Lys-719 of IkBζ does not participate in Lcn2 activation, which also appears to be supported by the finding that neither gene activation nor promoter association are prevented by replacement of the lysines with arginine, a residue that does not undergo acetylation (Figs. 4F and 5D).

Basic Residues in ANK7 of IkBζNS and Bcl-3 Are Involved in Association with Target DNA—Lys-717 and Lys-719 of IkBζNS are predicted to follow the second α helix in ANK7, and basic residues also exist at the corresponding sites in ANK7 of IkBζNS and Bcl-3 (Fig. 7A). As shown in Fig. 7B, association of IkBζNS with the promoter of the IkBζNS-regulated gene Il6 was prevented by simultaneous glutamate substitution for Lys-316, Arg-317, and Arg-319 in ANK7, although the substitution did not affect IkBζNS binding to p50. Similarly, glutamate substitution for Arg-354 and Lys-356 in ANK7 of Bcl-3 resulted in a loss of complex formation with the promoter of the Bcl-3-dependent gene Cxcl10 without affecting Bcl-3 interaction with p50 (Fig. 7C), and Bcl-3 (R354E/K356E) also failed to associate with the endogenous promoter of Cxcl10, as indicated by ChIP analysis (Fig. 7D). Thus, in IkBζNS and Bcl-3, basic residues that follow the second α helix in ANK7 appear to be involved in recognition of their target gene promoters. Furthermore, the R354E/K356E substitution in Bcl-3 resulted in loss of the p52-dependent Ccl2 activation in LPS-stimulated RAW264.7 cells (Fig. 7E), confirming the significance of the basic residues in ANK7.
The DNA-binding assays also revealed that the invariant asparagine residues in ANK1 of IkBζNS (Asp-60) (Fig. 7B) and Bcl-3 (Asp-127) (Fig. 7C and D) participate in association with their target gene promoters via direct binding to p50, similar to the corresponding residue of IkBζ (Asp-451) (Fig. 5A).

Activation of Lcn2 Involves Promoter Association with IkBζ via an Extra-κB Site in a Sequence-specific Manner—Activation of the mouse Lcn2 gene requires promoter association with the IkBζ-p50 complex via both the κB site (5'-GGGAATGTCCC-3' at positions -230 to -220 relative to the transcription start site) and its downstream region of 5'-CCCCTC-3' at positions -212 to -207 (23) (see Fig. 8A). To elucidate base specificity in the downstream sequence, we constructed a series of mutant Lcn2 promoters and tested their ability. Substitution of either guanine or adenine for cytosine at position -212 led to a loss of both interaction with IkBζ and IkBζ-mediated Lcn2 activation (Fig. 8B). On the other hand, they were only marginally impaired by thymine replacement at the corresponding position (Fig. 8B). Adenine but not thymine or guanine partially replaced cytosine at position -211 (Fig. 8C). At positions -210 (Fig. 8D) and -209 (Fig. 8E), cytosine was strictly required for both IkBζ binding to the Lcn2 promoter and IkBζ-mediated Lcn2 activation. By contrast, any of four bases fully functioned at positions -207 (Fig. 8F) and -209 (Fig. 8G). In the sequence CCCCTC at positions -212 to -207 of the mouse Lcn2 promoter, the preference for cytosine at positions -212 to -209 but not at position -208 or -207 (Fig. 8, B–G) appears to be consistent with the conservation of the first four cytosines but not the last two bases in the corresponding region from other mammals (Fig. 8A). These mutational analyses indicate that

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![Figure 6. Protein acetylation is not involved in IκBζ-mediated Lcn2 activation.](image)

A and B, effect of anacardic acid (A) and nicotinamide (B) on IκBζ-mediated Lcn2 activation. p50-/IκBζ-deficient MEFs were transfected with the following plasmids: the luciferase reporter plasmid pGL3-Basic containing the upstream region of Lcn2 (~1031/+54), the internal control plasmid pRL-TK, and the pcDNA3 vector or pcDNA3 for expression of FLAG-IκBζ and HA-p50. Cells were incubated for 12 h with the indicated concentrations of Lcn2-mediated activation. Results are means ± S.D. for three independent transfections. C, effect of the HDAC inhibitor TSA on IκBζ. HEK293T cells expressing FLAG-IκBζ were incubated for 6 h with 2 μM TSA. The cell lysates were applied to immunoprecipitation (IP) with anti-FLAG antibody. Precipitated FLAG-IκBζ as well as acid extracted histones in the nuclear extracts were analyzed by immunoblot with the indicated antibodies or by staining with CBB. Positions for marker proteins are indicated in kilodaltons.

IκBζ-responsive element (positions −212 to −209) requires the sequence 5′-YCCC-3′ (Y is pyrimidine). Taken together with the present findings, IκBζ activates the endogenous Lcn2 gene both via Asp-451-dependent direct interaction with p50 and Lys-717/Lys-719-involved association with the sequence 5′-YCCC-3′ downstream of the κB site.

Discussion

The nuclear IκB proteins IκBζ, Bcl-3, and IκBNS are thought to regulate NF-κB-dependent transcription by directly interacting with a p50 or p52 homodimer that binds to the κB site on target genes. However, the mechanism for formation of the regulatory complex has not been fully elucidated. In the present study, we show that IκBζ, comprising the N-terminal transactivation domain and the C-terminal ARD composed of seven ANK motifs, forms a transcriptionally active complex on its target gene Lcn2 both via Asp-451-mediated binding to p50 and via Lys-717/Lys-719-dependent interaction with the extra-κB site of the Lcn2 promoter. Asp-451 is present in the N-terminal region of ANK1, whereas the basic residues Lys-717 and Lys-719 exist in the C-terminal region of ANK7. We also demonstrate similar roles for both termini of the ARD in Bcl-3 and IκBNS, proposing a model for a common mechanism by which nuclear IκBs form a p50/p52-containing complex on target gene promoters.

Asp-451 in IκBζ ANK1 is strictly conserved during evolution (Fig. 1). Replacement of Asp-451 by threonine abrogates both association with a homodimer of the NF-κB subunit p50 (Fig. 1) and activation of Lcn2 via formation of the IκBζ-p50-DNA complex on the promoter (Figs. 3 and 5). The aspartate residue is also conserved among the nuclear IκBs, including Bcl-3 and IκBNS; however, it is replaced by threonine or serine in cytoplasmic IκBs such as IκBo and IκBβ (Fig. 1), which associate with a p50/p52 homodimer much less efficiently (13). The conservation among the nuclear IκBs is consistent with the present finding that the corresponding aspartate residues (Asp-127 in Bcl-3 and Asp-60 in IκBNS) are also crucial for interaction with a p50 or p52 homodimer (Figs. 1 and 2). It should be noted that Asp-451 in IκBζ is one of the very few residues that are completely conserved among nuclear IκBs but replaced in cytoplasmic IκBs. On the other hand, the presence of aspartate at this position by itself does not seem to be sufficient because a mutant IκBα carrying aspartate substitution for Thr-71 at the corresponding position of ANK1 as well as the wild-type protein fails to bind to a p50 homodimer (data not shown).

Although little is known about the tertiary structure of nuclear IκB-κB complexes, crystal structures of the cytoplasmic IκB proteins IκBo and IκBβ complexed with a p65-p50 heterodimer and a p65 homodimer, respectively, have been solved (45–47). If IκBζ interacts with NF-κB subunits in a manner similar to the cytoplasmic IκBs, then it seems likely that Asp-451 in IκBζ is positioned toward the C-terminally localized nuclear localization signal (NLS) in p50, a region that is required not only for nuclear localization of p50 but also for direct interaction of p50 with IκBζ (23, 48). Because of the dual role of the p50 NLS-containing region, the requirement of direct p50-IκBζ interaction for gene activation has not been established, although an NLS-truncated p50 protein did not activate IκBζ-dependent transcription, and thus IκBζ was assumed to interact with the promoter DNA via association with p50, which directly binds to the κB site (23, 48).

The conclusion appears to be strongly supported by the present findings that the D451T substitution in IκBζ abrogates interaction with p50 (Fig. 1), complex formation on both endogenous and exogenous Lcn2 promoter (Fig. 5), and activation of Lcn2 transcription (Fig. 3).

In addition to direct association with the κB-site-binding protein p50, IκBζ activates Lcn2 transcription by interacting with the extra-κB site of the Lcn2 promoter in which the basic residues Lys-717 and Lys-719 in the C-terminal region of IκBζ ANK7 likely play a major role (Figs. 4 and 5). The association does not appear to be mediated via direct binding to p50 because the K717E/K719E substitution in IκBζ leads to a loss of both promoter association and Lcn2 activation without affect-
ING direct contact of IκBζ with p50 (Figs. 4 and 5). In addition, the function of the lysine residues does not seem to require their modification, such as acetylation (Fig. 6), which is supported by the finding that arginine, a residue insusceptible to acetylation, fully serves in place of them (Figs. 4 and 5).

The extra-κB site required for IκBζ-dependent Lcn2 activation localizes seven bases downstream of the κB site in the Lcn2 promoter (23) and strictly requires the sequence 5′-YCCC-3′ as an IκBζ-responsive element, as shown in this study (Fig. 8). In both promoter association and Lcn2 activation, Lys-717 and Lys-719 in IκBζ are fully replaced by arginine residues (Figs. 4 and 5). It is known that lysine and arginine, but not glutamate, are often involved in direct interaction with DNA, which can be mediated not only via nonspecific interaction with the phos-
IκBζ Ankyrin Repeats Bind to NF-κB p50 and Promoter DNA

**FIGURE 8. Interaction of IκBζ with the Lcn2 promoter, and its activation requires the 5′-YCCC-3′ sequence downstream of the κB site.** A, the 5′ upstream promoter region of the mouse Lcn2 gene. The κB site and its downstream IκBζ-responsive element in the region are underlined. The nucleotide sequences of the corresponding region from various species are aligned. Mm, *Mus musculus*; Hs, *Homo sapiens*; Pa, *Papio anubis*; Rn, *Rattus norvegicus*. B–G, base preference of the extra-κB site in Lcn2 activation (top panels) and in formation of the three-species complex containing IκBζ, p50, and promoter DNA (bottom panels). For estimation of Lcn2 activation, p50/κB-deficient MEFs were transfected with the following plasmids: the luciferase reporter plasmid pGL3-Basic containing the upstream region of wild-type Lcn2 (~2301-220); a mutant Lcn2 with the indicated base replacement, or the IκBζ-independent gene Sele (~445/+105); the internal control plasmid pRL-TK, and pcDNA3 for expression of FLAG-IκBζ and HA-p50. Luciferase activities were determined as described under Experimental Procedures. Each graph represents the mean ± S.D. obtained from three independent transfections. For estimation of protein-DNA complex formation, GST-IκBζ was incubated with His-p50 in the presence of the DNA fragment of wild-type Lcn2 (~500/+50), a mutant Lcn2 with the indicated base replacement, or Sele (~445/+105). After the complex was pulled down with glutathione-Sepharose-4B beads, the co-precipitated DNA was amplified by PCR, and the product was analyzed by agarose gel electrophoresis.

phosphate moiety of DNA but also via recognition of a DNA base, especially a guanine (39, 40). In this context, it should be noted that the IκBζ-responsive element (5′-YCCC-3′) of the Lcn2 promoter is abundant in the Lys/Arg-recognizing base guanine in the antisense strand. The presence of multiple guanines in the element may be in agreement with the present conclusion that Lys-717 and Lys-719 each contribute to element recognition because single glutamate substitution for either lysine residue results in only partial loss of the activity of IκBζ (Fig. 4). Correct orientation of the side chains of Lys-717 and Lys-719 toward target DNA also seems to be important for element recognition, as indicated by the finding that replacement of the inflexible residue Gly-718, which is the intervening amino acid between the lysines and is strictly conserved during evolution, by the inflexible residue proline abrogates both binding to the Lcn2 promoter and transcription of Lcn2 (Fig. 5).

The NF-κB family proteins serve as a homo- or heterodimer to bind to a κB DNA response element in the promoters of distinct inducible genes, thereby playing their respective roles in gene regulation (1–4). The transcriptional specificity of NF-κB dimers is generally thought to be coded within the κB site sequences (27, 49). Indeed, in the case of the Lcn2 promoter, its κB site effectively interacts with the NF-κB p50 homodimer but not with the p52 homodimer (Fig. 3), although both homodimers are capable of binding to IκBζ (Fig. 2). As a result, in contrast to p50, p52 fails to induce IκBζ-dependent activation of the Lcn2 gene (Fig. 3). In addition to the κB site sequence itself, the ability of NF-κB dimers to function on a particular promoter is also considered to be determined by extra-κB sites recognized via co-activator proteins (16, 50). However, such a co-activator was not previously identified. This study demonstrates that IκBζ, a co-activator of p50, activates the Lcn2 gene not only via binding to p50 but also by associating with the IκBζ-responsive element (5′-YCCC-3′) downstream of the κB site, providing evidence that an extra-κB site and its interacting co-activator do determine the transcriptional specificity of NF-κB dimers.

Bcl-3 and IκBζ are the putative second and third examples that determine the transcriptional specificity of NF-κB dimers via association with extra-κB sites. These nuclear IκB proteins, capable of binding to a p50 and p52 homodimer (Figs. 1 and 2), have conserved basic residues that follow the second α helix in the C-terminal region of ANK7; the residues appear to be involved in association of Bcl-3 and IκBζ with promoters of their target genes (Fig. 7). The crucial role for Arg-354 and Lys-356 of Bcl-3 and IκBζ in promoter binding and gene activation agrees well with a model in which the C-terminal region of ANK7; the residues appear to be involved in association of Bcl-3 and IκBζ with promoters of their target genes (Fig. 7). The crucial role for Arg-354 and Lys-356 of Bcl-3 and IκBζ in promoter binding and gene activation agrees well with a model in which the C-terminal region of ANK7; the residues appear to be involved in association of Bcl-3 and IκBζ with promoters of their target genes (Fig. 7). The crucial role for Arg-354 and Lys-356 of Bcl-3 and IκBζ in promoter binding and gene activation agrees well with a model in which the C-terminal region of ANK7; the residues appear to be involved in association of Bcl-3 and IκBζ with promoters of their target genes (Fig. 7).
region of ANK1 and interaction with an extra-κ site via recognition by the conserved lysine/arginine residues that locate C-terminally to the second α helix of ANK7.

**Experimental Procedures**

**Cells, Antibodies, and Reagents—**MEFs doubly deficient in NF-κB p50 and IkBζ (Nfkbia−/−;Nfkbiz−/−) were prepared as described previously (23). Mouse BMMs were obtained as described previously (22, 24). All animals were housed and maintained in a specific pathogen-free animal facility at Kyushu University. All experiments were performed in strict accordance with the guidelines for proper conduct of animal experiments (Science Council of Japan). The experimental protocol was approved by the Animal Care and Use Committee of Kyushu University (permit numbers: A24-042 and A26-102).

All efforts were made to minimize the numbers of animals and their suffering. MEFs, BMMs, HEK293T cells, and mouse macrophage-like RAW 264.7 cells were maintained in Dulbecco's modified Eagle's medium supplemented with 10% heat-inactivated fetal bovine serum, penicillin (100 units/ml), and streptomycin (100 μg/ml) at 37 °C under 5% CO₂.

Anti-FLAG (M2) (catalog no. F3165) and anti-β-tubulin (TUB 2.1, catalog no. T4026) mouse monoclonal antibodies and anti-FLAG rabbit polyclonal antibodies (catalog no. F7425) were purchased from Sigma-Aldrich, an anti-HA rat monoclonal antibody (3F10) (catalog no. 11 867 431 001) from Roche Applied Science, an anti-His₅ monoclone antibody (catalog no. 34660) from Qiagen, an anti-RNA polymerase II rabbit polyclonal antibody (catalog no. sc-47778) and anti-RNA polymerase II rabbit polyclonal antibody (catalog no. sc-899) from Santa Cruz Biotechnology, and an anti-acetylated lysine rabbit polyclonal antibody (catalog no. 9441) from Cell Signaling Technology. Anti-IkBζ rabbit polyclonal antibodies were raised against IkBζ (1–100) and prepared as described previously (51). TSA and anacardic acid were purchased from Calbiochem and nico
tinamide from Sigma-Aldrich

**Plasmid Construction—**The mouse cDNAs encoding full-length IkBζ (amino acid residues 1–728), IkBζ-(414–728), p50 of 366 amino acid residues, and p65 of 325 amino acids were prepared as described previously (9, 33). The cDNA for mouse full-length IkBNS (amino acid residues 1–327) were prepared by RT-PCR using mRNA from mouse BMMs stimulated with LPS (List Biological Laboratories). The cDNAs for mouse full-length Bcl-3 (amino acid residues 1–448) and p52 of 415 amino acid residues were prepared by RT-PCR using mRNA from LPS-stimulated RAW 264.7 cells. The cDNA fragments encoding Bcl-3 (122–362) and p52 (1–341) for bacterial expression were prepared by PCR using their respective full-length cDNAs. Mutations leading to the indicated amino acid substitutions or truncations were introduced by PCR-mediated site-directed mutagenesis. The cDNA fragments were ligated to the following expression vectors: pGEX-6P-2 (GE Healthcare) and pMALc2 (New England Biolabs) for bacterial expression of proteins fused to GST and maltose-binding protein (MBP), respectively; pRSFDuet-1 (Novagen) modified for expression of hexahistidine (His)-tagged proteins in Escherichia coli (52); pcDNA3 (Invitrogen) for expression of FLAG- or HA-tagged proteins in mammalian cells; and pBABE-puro (Cell Biolabs) for retroviral transduction in BMMs. The 5’ upstream region (−1031/+54 or −500/+50) of the lipocalin-2-encoding gene Lcn2 or the 5’ upstream region (−445/+105) of the E-selectin-encoding gene Sele was ligated to the luciferase reporter plasmid pGL3-Basic (Promega) as described previously (23). The 5’ upstream regions of the following genes were amplified by PCR using mouse genomic DNA as a template and subcloned into pGL3-Basic: the IL-6-encoding gene Il6 (−1217/+50), the IP-10-encoding gene Cxcl10 (−500/+50), and the MCP-1-encoding gene Ccl2 (−2777/+76). Mutations in the κ site of the following gene promoters were introduced by PCR-mediated site-directed mutagenesis: Lcn2 (−230/−220), 5’T-CCGAAG-TGTCCCT-3’ to 5’T-TATAAATGTTAAT-3’ (23, 24); Il6 (−73/−64), 5’T-GGGATTTTCCA-3’ to 5’T-TATATTTTAAA-3’; and Cxcl10 (−113/−103), 5’T-AGGGGACTTTCC-3’ to 5’T-ATAATACCTTAAT-3’ (note that mutated nucleotides are underlined). All of the constructs were sequenced for confirmation of their identity.

**GST Pulldown Assay—**GST- and His-tagged proteins were purified as described previously (52, 53) and incubated for 20 min at 4 °C in 500 μl of buffer A (137 mM NaCl, 2.7 mM KCl, 0.5% Triton X-100, 1 mM DTT, 8.1 mM Na₂HPO₄, and 1.5 mM KH₂PO₄ (pH 7.4)). A slurry of glutathione-Sepharose 4B beads (GE Healthcare) was subsequently added, followed by further incubation for 40 min at 4 °C. After washing three times with the buffer above, the proteins were eluted from the beads with 20 mM glutathione in 150 mM NaCl, 2 mM DTT, and 100 mM Tris-HCl (pH 8.8). The eluate was subjected to SDS-PAGE, followed by staining with Coomassie Brilliant Blue (CBB).

**Immunoprecipitation Analysis—**HEK293T cells were transfected with the indicated expression plasmids using X-tremeGENE HP DNA Transfection Reagent (Roche Applied Science) and cultured for 24 or 48 h. Cells were lysed by sonication at 4 °C in 500 μl of lysis buffer (137 mM NaCl, 2.7 mM KCl, 1% Nonidet P-40, 8.1 mM Na₂HPO₄, and 1.5 mM KH₂PO₄ (pH 7.4)) supplemented with Complete™ Protease Inhibitor Cocktail (Roche Applied Science). Proteins in the cell lysate were immunoprecipitated using anti-FLAG antibody (M2) and protein G-Sepharose (GE Healthcare). The precipitants were analyzed by immunoblot with the anti-FLAG rabbit polyclonal antibody or the anti-HA (3F10) rat monoclonal antibody. The blots were developed using ECL-Prime (GE Healthcare) for visualization of the antibodies.

**Detection of Acetylated Proteins—**FLAG-IkBζ expressed in HEK293T cells was incubated for 6 h with 2 μM TSA and immunoprecipitated from the cell lysates as described above, with the exception that 2 μM TSA and 20 mM nicotinamide were used during immunoprecipitation to prevent deacetylation. The precipitated proteins were applied to immunoblot analysis with anti-acetylated lysine rabbit polyclonal antibody. Histones were prepared from HEK293T cells by acid extraction as described by Nightingale et al. (54). For detection of acetylated histones, the acid extracts were subsequently analyzed by immunoblot with the anti-acetylated lysine rabbit polyclonal antibody.

**Luciferase Reporter Assay—**Nfkbia−/−;Nfkbiz−/− MEFs or RAW264.7 cells were transfected with the luciferase reporter plasmid pGL3-Basic containing the indicated promoter, the
internal control plasmid pRL-TK (Promega), and the indicated plasmids for protein expression using X-tremeGENE HP DNA Transfection Reagent. The luciferase activities were determined by the Dual-Luciferase® reporter assay system (Promega). For estimation of protein levels, cell lysates were analyzed by immunoblot with anti-FLAG (M2), anti-HA, anti-β-tubulin, or anti-β-actin antibody.

Analysis of Protein-DNA Interaction—Analysis of protein-DNA interaction was performed as described previously (23). GST-Ikbζ, GST-IkbNS, or MBP-Bcl-3 was incubated with His-p50 for 20 min at 4 °C in 500 μl of buffer A containing salmon sperm DNA (0.1 mg/ml), or His-p50, His-p52, or His-p65 alone was incubated for 20 min at 4 °C in 500 μl of buffer B (150 mM NaCl, 5% glycerol, 1 mM DTT, 0.5% Triton X-100, and 25 mM Tris-HCl (pH 8.0)) containing salmon sperm DNA (0.1 mg/ml). A slurry of glutathione-Sepharose 4B, Amylose Resin (New England Biolabs), or COSMOGEL® His-Accept (Nacalai Tesque) was added in a GST, MBP, or His pulldown assay, respectively. The indicated DNA fragment (0.1 pmol) was subsequently added to the mixture, followed by further incubation for 40 min at 4 °C. After washing with buffer A for a GST or MBP pulldown assay or with buffer B containing 25 mM imidazole for a His pulldown assay, the protein-DNA complex was eluted from glutathione-Sepharose 4B with an elution buffer (20 mM glutathione, 150 mM NaCl, 2 mM DTT, and 100 mM Tris-HCl (pH 8.8)), from amylose resin with buffer A containing 20 mM maltose, or from COSMOGEL® His-Accept with buffer B containing 1 mM imidazole. The eluted protein was applied to SDS-PAGE, followed by CBB staining or immunoblot analysis with anti-FLAG (M2), anti-HA, anti-β-tubulin, or anti-β-actin antibody.

Analysis of protein–DNA interaction was performed as described previously (23). GST-Ikbζ, GST-IkbNS, or MBP-Bcl-3 was incubated with His-p50 for 20 min at 4 °C in 500 μl of buffer A containing salmon sperm DNA (0.1 mg/ml), or His-p50, His-p52, or His-p65 alone was incubated for 20 min at 4 °C in 500 μl of buffer B (150 mM NaCl, 5% glycerol, 1 mM DTT, 0.5% Triton X-100, and 25 mM Tris-HCl (pH 8.0)) containing salmon sperm DNA (0.1 mg/ml). A slurry of glutathione-Sepharose 4B, Amylose Resin (New England Biolabs), or COSMOGEL® His-Accept (Nacalai Tesque) was added in a GST, MBP, or His pulldown assay, respectively. The indicated DNA fragment (0.1 pmol) was subsequently added to the mixture, followed by further incubation for 40 min at 4 °C. After washing with buffer A for a GST or MBP pulldown assay or with buffer B containing 25 mM imidazole for a His pulldown assay, the protein-DNA complex was eluted from glutathione-Sepharose 4B with an elution buffer (20 mM glutathione, 150 mM NaCl, 2 mM DTT, and 100 mM Tris-HCl (pH 8.8)), from amylose resin with buffer A containing 20 mM maltose, or from COSMOGEL® His-Accept with buffer B containing 1 mM imidazole. The eluted protein was applied to SDS-PAGE, followed by CBB staining or immunoblot analysis with anti-FLAG (M2), anti-HA, anti-β-tubulin, or anti-β-actin antibody.

Activation of the Endogenous Lcn2 Gene by Ikbζ—Retroviral expression of wild-type Ikbζ or a mutant protein with the D451T or K717E/K719E substitution was performed according to the method of He et al. (55). Briefly, HEK293T cells were transfected with the Moloney Murine Leukemia virus-ΨE helper plasmid and pBABE-puro-3×FLAG-Ikbζ, and the culture supernatant containing the retrovirus was collected. Ikbζ-deficient BMMs were infected with the retrovirus and cultured for 2 days. The cells were treated for 4, 8, or 12 h with or without LPS (100 ng/ml). Expression of Ikbζ was detected by immunoblot with the anti-Ikbζ antibody. The mRNA product of the endogenous Lcn2 gene was estimated by quantitative real-time RT-PCR as described previously (56). Briefly, total RNAs were extracted using TRIzol (BIOLINE) according to the instructions of the manufacturer, and 1 μg of the RNA was reverse-transcribed by ReverTra Ace® reverse transcriptase (TOYOBO) using an oligo(dT) primer, followed by real-time PCR using SYBR® premix Ex Taq™ (Takara Bio) on the Roter-Gene 6200 system (Corbett). The primer pairs used were 5′-AACGGACGTGCCTCGATT-3′ and 5′-GGTTGGGACAGAAGATGA-3′ for Lcn2 and 5′-AAACGGACTGCGGCAAC-3′ and 5′-TAAACGGATGTTGGGACATG-3′ for the control gene Rpl32.

ChiP Analysis—RAW264.7 cells expressing Ikbζ or Bcl-3 under the control of the sheep metallothionein la promoter (−600/+72) were prepared as described previously (22). The stable transformants were incubated in the presence of 50 μM ZnSO4 and LPS (100 ng/ml), and ChiP analysis was performed according to a method described previously (24, 56). Cells were fixed for 10 min at 25 °C with 1% formaldehyde and washed with ice-cold phosphate-buffered saline. After sonication, a chromatin-containing solution was precleared with Protein G-Sepharose 4 Fast Flow (GE Healthcare), followed by incubation overnight at 4 °C with the indicated antibody. Antigen-antibody complexes on the resin were washed sequentially with a low-salt wash buffer (150 mM NaCl, 2 mM EDTA, 0.1% SDS, 1% Triton X-100, and 20 mM Tris-HCl (pH 8.0)), a high-salt wash buffer (500 mM NaCl, 2 mM EDTA, 0.1% SDS, 1% Triton X-100, and 20 mM Tris-HCl (pH 8.0)); a LiCl wash buffer (250 mM LiCl, 1 mM EDTA, 1% sodium deoxycholate, 1% Nonidet-P40, and 10 mM Tris-HCl (pH 8.0)), and TE (1 mM EDTA and 10 mM Tris-HCl (pH 8.0)). DNA-protein complexes were eluted with elution buffer containing 1% SDS and 100 mM sodium bicarbonate. After cross-links were reversed by overnight incubation at 65 °C, Proteinase K (0.1 mg/ml) was added and incubated for 3 h at 45 °C. Purified DNA was subjected to PCR to detect the following regions using specific primers as follows: the Lcn2 promoter region containing the CCCCCC element (−212/−207), 5′-CCCCCTGTGCCCCCTGCAGC-3′, and 5′-TCTGTTGAATACCTTGGCGAAGAT-3′ and the Cxcl10 promoter region, 5′-TCAAGGTCATGGTTGCTACAAGA-3′ and 5′-GGGAAGTCCCCCTGAAACCGA-3′.

Secondary Structure Prediction—The secondary structure of the ANK7 in Ikbζ and IkbNS was predicted using the server-side program PSIPRED (57, 58).

Author Contributions—A. K., S. Y., and H. S. designed the study. A. K. performed the experiments. A. K., S. Y., and H. S. analyzed the data and wrote the paper. All authors analyzed the results and approved the final version of the manuscript.

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