AMAP1 as a negative-feedback regulator of nuclear factor-κB under inflammatory conditions

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NF-κB is a major transcriptional factor regulating many cellular functions including inflammation; therefore, its appropriate control is of high importance. The detailed mechanism of its activation has been well characterized, but that of negative regulation is poorly understood. In this study, we showed AMAP1, an Arf-GTPase activating protein, as a negative feedback regulator for NF-κB by binding with IKKβ, an essential kinase in NF-κB signaling. Proteomics analysis identified AMAP1 as a binding protein with IKKβ. Overexpression of AMAP1 suppressed NF-κB activity by interfering the binding of IKKβ and NEMO, and deletion of AMAP1 augmented NF-κB activity. The activation of NF-κB induced translocation of AMAP1 to cytoplasm from cell membrane and nucleus, which resulted in augmented interaction of AMAP1 and IKKβ. These results demonstrated a novel role of AMAP1 as a negative feedback regulator of NF-κB, and presented it as a possible target for anti-inflammatory treatments.

Nuclear Factor-κB (NF-κB) consists of a family of transcription factors (p65 or RelA, p50, p52, c-Rel, and RelB) that play critical roles in inflammation, immunity, cell proliferation, differentiation and survival. It generally exists as a homo- or heterodimer in the cytosol that is bound to the inhibitor of κB (IκB) (see review). In response to a wide variety of stimuli, including inflammatory cytokines, IκB is phosphorylated and degraded via the ubiquitin pathway, which is followed by NF-κB translocation to the nucleus and activation of transcription. Serine phosphorylation of IκB is mediated by a large multi-unit complex containing two catalytic subunits (IKKα and IKKβ) and the regulatory subunit IKKγ or NEMO. Among these subunits, IKKβ is the essential kinase mediating IκB phosphorylation in most cell types, and germline deletion of IKKβ results in embryonic lethality due to massive liver apoptosis, similar to the phenotype of p65-knockout mice.

NF-κB signaling proteins including these kinases are essential for maintaining living body as shown by IKKβ or p65 knockout mouse with phenotype of embryonic lethality, but they also have been implicated in the pathogenesis of many human diseases, including cancer. Therefore, controlling their activities appropriately is of high importance. Although the mechanism for its activation was been well analyzed as described above, that for negative regulation is poorly explored.

We recently reported that IKKβ has a kinase-independent role in regulating widespread cellular responses and cell signaling, implicating that there are still undiscovered roles of IKKβ. To explore the roles more extensively, we identified A Multiple-domain Arf-GAP Protein 1 (AMAP1) as a binding partner of IKKβ by proteomic analyses. AMAP1, also called ASAP1 or DDEF1, is an Arf-GTPase-activating protein that functions on membrane surfaces to catalyze the hydrolysis of GTP bound to Arf, and plays major roles in the regulation of membrane remodeling and cytoskeletal organization, cellular migration and tumor invasion and metastasis. More importantly, clinical studies have indicated that AMAP1 is dramatically up-regulated in advanced cancers. The roles of AMAP1 in the inflammatory responses are of high interest considering the close relationship between cancer and inflammation; however, the roles have been poorly understood. Interestingly, Haque et al.
reported the inhibitory effect of AMAP1 (ASAP1) on the production of proinflammatory mediators\(^2\), but the detailed mechanism remains unknown.

In this report, we carefully examined the involvement of AMAP1 in the regulation of NF-\(\kappa\)B, and found that it has an unanticipated role to interfere the IKK\(\beta\)–NEMO binding, which results in negative-feedback regulation of NF-\(\kappa\)B activity. This report describes the detailed role of AMAP1 in the inflammatory responses, and AMAP1 can be a new target for developing anti-inflammatory treatments.

**Results**

**AMAP1 interacts with IKK\(\beta\).** To discover novel roles of IKK\(\beta\), we started with identifying proteins that bind to IKK\(\beta\). Three kinds of FLAG-tagged IKK\(\beta\) were overexpressed in HEK293T cells, and binding proteins were immunoprecipitated using anti-FLAG antibody. The samples were subjected to proteomic analyses using the nano-LC/MALDI-TOF system. Of the proteins identified repeatedly, we focused on AMAP1 since it is clinically important due to its augmented expression in cancer\(^15,16,19,22,24\), which is one of the major inflammatory diseases\(^29\). To confirm the physical association of IKK\(\beta\) and AMAP1, we performed co-immunoprecipitation (Co-IP) assays using the lysate of HEK293T cells co-overexpressed with HA-tagged AMAP1 and FLAG-tagged IKK\(\beta\). IKK\(\beta\) and AMAP1 were detected in the precipitates with either the anti-HA antibody or the anti-FLAG antibody, respectively (Fig. 1a).

Next, we determined the binding sites of these proteins. Based on the functional modules of IKK\(\beta\), including an amino-terminal kinase domain (KD), a ubiquitin-like domain (ULD) and a carboxyl-terminal NEMO Binding Domain (NBD)\(^2\), we divided IKK\(\beta\) into 4 overlapping FLAG-tagged fragments. The constructs are IKK\(\beta\)-A1 comprising the KD, ULD and SDD, IKK\(\beta\)-A2 comprising the KD and ULD, IKK\(\beta\)-A3 comprising only the KD, and IKK\(\beta\)-A4 comprising the ULD, SDD and NBD (Fig. 1b), and each construct was expressed in HEK293T cells along with HA-tagged AMAP1. We found that AMAP1 was co-precipitated with IKK\(\beta\)-A4, but not with any other construct of IKK\(\beta\) (Fig. 1c), indicating that AMAP1 binds to IKK\(\beta\) at the C-terminal region containing the NBD. Further, we examined which region of AMAP1 is essential for its binding to IKK\(\beta\). Each of 6 GST-tagged fragments of AMAP1, which are Bar (Bin-Amphiphysin-Rvs), PH (Pleckstrin Homology), Arf-GAP, ANK (Ankyrin repeat), PRD (Proline Rich Domain) and SH3 (Src Homology3) domains (Fig. 1d), was expressed in HEK293T cells along with FLAG-tagged IKK\(\beta\). We found only SH3 domain construct was co-precipitated with IKK\(\beta\) as shown in Fig. 1e. To confirm it further, we made construct of SH3-domain deleted AMAP1 (AMAP1\(\Delta\)SH3). Cells were expressed with full-length AMAP1 or AMAP1\(\Delta\)SH3 along with FLAG-tagged IKK\(\beta\), and we found only full-length AMAP1, not AMAP1\(\Delta\)SH3, was co-precipitated with IKK\(\beta\) (Supplementary Fig. S1 online). Taken together, these results indicated that the SH3 domain of AMAP1 is responsible for the binding with IKK\(\beta\), which implicates that the GTPase activity of AMAP1 is not involved in the interaction with IKK\(\beta\).

**AMAP1 negatively regulates NF-\(\kappa\)B.** The C-terminal region of IKK\(\beta\) identified as the binding site for AMAP1 contains the binding domain for NEMO/IKK\(\gamma\), which is the regulatory subunit of the IKK complex\(^3\). It prompted us to verify the effect of AMAP1 on the association of IKK\(\beta\) and NEMO because it is required for NF-\(\kappa\)B activation\(^26\). As shown in Fig. 2a, overexpression of AMAP1 inhibited the binding of IKK\(\beta\) and NEMO, and it was in a dose-dependent manner (Supplementary Fig. S2 online). In addition, the increased level of NEMO could mitigate the interaction between AMAP1 and IKK\(\beta\) (Fig. 2b). Taken together, there is competition between AMAP1 and NEMO in IKK\(\beta\) interaction. As expected from these results, overexpression of AMAP1 suppressed NF-\(\kappa\)B activity (Fig. 2c), and it was confirmed by less translocation of p65 into nucleus after IL-1\(\beta\) stimulation in AMAP1-overexpressed cells (Fig. 2d). Additionally, suppression of AMAP1 by shRNA enhanced NF-\(\kappa\)B activity (Fig. 2e) and augmented p65 translocation into the nucleus after IL-1\(\beta\) stimulation (Fig. 2f). Thus, AMAP1 interferes the binding of IKK\(\beta\) and NEMO, and negatively regulates NF-\(\kappa\)B.

**NF-\(\kappa\)B activation induces AMAP1 translocation to cytoplasm and augments the AMAP1-IKK\(\beta\) interaction.** In the experiments mentioned above, we sometimes observed that more amount of AMAP1 was detected after treatment with IL-1\(\beta\) (Supplementary Fig. S3a online). This observation coincides with previous report showing LPS significantly boosted AMAP1 protein levels within 15 min\(^2\). However, considering that 15 min is too short for synthesis or degradation of proteins, it was highly possible that AMAP1 might not be completely dissolved in cell lysis buffer because AMAP1 can be localized in lipid rafts\(^11-14\) and nucleus\(^27\), both of which might not be fully dissolved by 1% Triton X-100 of cell lysis buffer used in this study. Indeed, we also observed similar change of AMAP1 by LPS stimulation, but it disappeared when cells were lysed in 1% SDS cell lysis buffer (Supplementary Fig. S3b online). Therefore, we hypothesized that NF-\(\kappa\)B activation induces AMAP1 translocation to the cytoplasm from cell membrane or nucleus, which could result in detection of more AMAP1 in cell lysate by 1% Triton X-100. As shown in Fig. 3a, fractionation by sucrose gradient indicated that IL-1\(\beta\) stimulation induced the translocation of AMAP1 from lipid rafts to the cytoplasm. We also performed confocal immunocytochemistry with using human umbilical ven endothelial cells (HUVECs) since HEK293T cells were not appropriate for observation of intracellular localization due to their large nuclei and thin cytoplasm. It was hard to see AMAP1 staining on cellular membrane, but we observed that IL-1\(\beta\) stimulation induced translocation of AMAP1 in nuclei to the cytoplasm (Fig. 3b). Finally, we repeated the sucrose gradient fractionation with HUVECs, and observed the translocation of AMAP1 from the membrane in response to IL-1\(\beta\) stimulation (Fig. 3c), similarly with HEK293T cells.

Considering that IKK\(\beta\) is nearly entirely cytoplasmic protein\(^26,29\), the translocation of AMAP1 to cytoplasm in response to IL-1\(\beta\) stimulation raised the question as to whether the activation of NF-\(\kappa\)B would strengthen the interaction between AMAP1 and IKK\(\beta\). As shown in Fig. 3d and e, more IKK\(\beta\) was co-precipitated with anti-AMAP1 antibody after IL-1\(\beta\) stimulation, and conversely, more AMAP1 was co-precipitated with anti-IKK\(\beta\) antibody after IL-1\(\beta\) stimulation, revealing that NF-\(\kappa\)B activation enhances the interaction between AMAP1 and IKK\(\beta\).

**Discussion**

Most biological processes require positive and negative regulatory mechanisms to maintain equilibrium. The results from this study suggest a novel mechanism for NF-\(\kappa\)B termination and contribute new knowledge to the mechanisms of NF-\(\kappa\)B regulation. So far, the re-synthesis of IkB proteins induced by the activation of NF-\(\kappa\)B is a widely accepted mechanism for terminating the NF-\(\kappa\)B response. These proteins enter the nucleus, remove NF-\(\kappa\)B from the DNA and re-localize it to the cytosol\(^40-33\). Here, we show that NF-\(\kappa\)B activation induces AMAP1 translocation to cytoplasm from nucleus or lipid rafts and accelerates binding of IKK\(\beta\) and AMAP1, which inhibits the association of IKK\(\beta\) and NEMO, eventually leading to inactivation of NF-\(\kappa\)B. Taken together, AMAP1 works as an inhibitory feedback mechanism for regulating the NF-\(\kappa\)B pathway (Fig. 4).

Recent studies have established strong support for the critical role of NF-\(\kappa\)B in cancer. Activation of NF-\(\kappa\)B controls multiple cellular processes in cancer, including inflammation, transformation, proliferation, angiogenesis, invasion, metastasis, chemoresistance and...
Figure 1 | AMAP1 interacts with IKKβ in vivo. (a) The binding of AMAP1 and IKKβ. FLAG-tagged IKKβ and HA-tagged AMAP1 were expressed in HEK293T cells and immunoprecipitated with anti-FLAG and anti-HA antibodies. AMAP1 and IKKβ were detected in the precipitate by immunoblotting with anti-FLAG and anti-HA antibodies. (b) Schematic representation of IKKβ and its mutants. (c) FLAG-tagged IKKβ/mutants and HA-tagged AMAP1 were expressed in HEK293T cells and immunoprecipitated with anti-FLAG antibody. AMAP1 was detected in the precipitate by immunoblotting with anti-HA antibody. (d) Schematic representation of AMAP1 and its mutants. (e) FLAG-tagged IKKβ and GST-tagged AMAP1/mutants were expressed in HEK293T cells and immunoprecipitated with anti-FLAG antibody. AMAP1 and its mutants were detected in the precipitate by immunoblotting with anti-GST antibody. The data shown are from one representative experiment of the three that were performed. Full-length blots are presented in Supplementary Figure 4.
Figure 2 | AMAP1 negatively regulates NF-κB. (a) AMAP1 interfered with the association of IKKβ and NEMO. FLAG-tagged IKKβ and NEMO were expressed in HEK293T cells along with control vector or HA-tagged AMAP1 and immunoprecipitated with anti-FLAG antibody. AMAP1, IKKβ and NEMO were detected in the precipitates by immunoblotting with anti-FLAG, anti-HA and anti-NEMO antibodies. (b) The induction of NEMO reduced the interaction of IKKβ and AMAP1. FLAG-tagged IKKβ and HA-tagged AMAP1 were expressed in HEK293T cells along with control or NEMO vector and immunoprecipitated with anti-FLAG antibody. AMAP1, IKKβ and NEMO were detected in the precipitates by immunoblotting with anti-FLAG, anti-HA and anti-NEMO antibodies. (c) NF-κB activity from nuclear extracts of overexpressed AMAP1 and control cells with or without IL-1β (2.5 ng/mL) treatment for 30 min. (d) Immunoblots of AMAP1 (whole cell) and p65 (nuclear extract) of HEK293T cells transfected with pcDNA3.1-HA-AMAP1 (AMAP1 overexpression) or pcDNA3.1 (control) and the cumulative, quantitative densitometry data. (e) NF-κB activity from nuclear extracts of shAMAP1 and shControl cells with or without IL-1β (2.5 ng/mL) treatment for 30 min. (f) Immunoblots of AMAP1 (whole cell) and p65 (nuclear extract) of HEK293T cells transfected with AMAP1 shRNA (shAMAP1) or Control shRNA (shControl) with or without IL-1β (2.5 ng/mL) and the cumulative, quantitative densitometry data. The data from three independent experiments are shown. Error bar: ±SD. * P < 0.05, ** P < 0.01 in a two-sided, Student’s t-test. Full-length blots are presented in Supplementary Figure 4.
Figure 3 | IL-1β induces AMAP1 translocation and the AMAP1 and IKKβ interaction. (a) Representative AMAP1 immunoblots of fractions from sucrose gradient centrifugation of HEK293T cells treated with or without IL-1β (2.5 ng/mL) for 30 min and the cumulative, quantitative densitometry data from three independent experiments are shown. (b) HUVECs were treated with or without IL-1β (2.5 ng/mL) for 30 min and then stained with anti-AMAP1 antibody, WGA and DAPI (n = 3). Scale bar = 50 μm. (c) Representative AMAP1 immunoblots of fractions from sucrose gradient centrifugation of HUVECs treated with or without IL-1β (2.5 ng/mL) for 30 min and the cumulative, quantitative densitometry data from three independent experiments are shown. Representative blots from three immunoprecipitation experiments using anti-AMAP1 (d) and anti-IKKβ (e) antibodies in HUVECs treated with or without IL-1β (2.5 ng/mL) for 30 min. The graph represents the cumulative, quantitative densitometry data of the IKKβ blot in anti-AMAP1 antibody precipitates (d), of the AMAP1 blot in anti-IKKβ antibody precipitates (e). Error bar: ±SD. * P < 0.05, *** P < 0.005 in a two-sided, Student’s t-test. Full-length blots are presented in Supplementary Figure 4.
radioresistance. Considering the role of AMAP1 explored in this study, augmented expression of AMAP1 observed in cancer might indicate overwhelming NF-kB activation and self-repairing against cancer.

In summary, this study rediscovered AMAP1 as a negative regulator of NF-kB activity in a feedback mechanism, and AMAP1 could be a novel target for treatment or prevention against inflammatory diseases including cancer.

Methods

Reagents. The mammalian expression vectors that were used included pcDNA3.1-HA-AMAP1 and pcDNA3.1 as a control for AMAP1 overexpression, a series of mammalian expression vectors for IKKβ and its mutants (pRES-AcGFPIKK2-WT, pRES-AcGFPIKK2-A1, pRES-AcGFPIKK2-A2, pRES-AcGFPIKK2-A3 and pRES-AcGFPIKK2-A4) and a series of mammalian expression vectors for AMAP1 and its mutants (pEBG-AMAP1-FL, pEBG/AMAP1-Bar, pEBG/AMAP1-PH, pEBG/AMAP1-AirGAP, pEBG/AMAP1-AN, pEBG/AMAP1-PRD and pEBG/AMAP1-SH3). The pcDNA3-HA-human NEMO plasmid (Addgene plasmid 13512) to express NEMO was obtained from Professor Kunliang Guan (University of California, San Diego, USA). Small hairpin RNA (shRNA) expression constructs for silencing AMAP1 (shAMAP1) and control shRNA (shControl) were purchased from GeneCopeia, Inc. (Rockville, MD, USA).

The antibodies that were used included anti-IKKβ (Millipore, Billerica, MA, USA); anti-AMAP1 (named anti-DDF1, Santa Cruz Biotechnology, Dallas, Texas, USA); anti-HA (Sigma-Aldrich, St. Louis, MO, USA); monoclonal ANTI-FLAG™ M2 (Sigma-Aldrich); a series of antibodies to anti-IKKβ, anti-phospho-IκBα, anti-IκBα, anti-p65, anti-IκBα, anti-GAPDH, anti-TATA box binding protein, anti-vimentin, anti-mouse IgG HRP-linked antibody and anti-rabbit IgG HRP-linked antibody (Cell Signaling Technology, Boston, MA, USA (CST)); anti-mouse IgG HRP and anti-rabbit IgG HRP (e Bioscience, Inc. San Diego, CA, USA); Alexa Fluor 488 goat anti-rabbit IgG and Alexa Fluor 594 goat anti-mouse IgG (Invitrogen, Foster City, CA, USA); and Alexa Fluor 488 WGA. The reagents that were used included interleukin-1β (Wako Pure Chemical Industries, Ltd., Osaka, Japan (Wako)), LPS (Sigma-Aldrich) and NE-PER Nuclear and Cytoplasmic Extraction Reagents (Thermo Scientific, Rockford, IL, USA).

Cell culture. The human embryonic kidney 293T cell line (HEK293T) was maintained in DMEM (Wako) containing 10% FBS (Gibco, Foster City, CA, USA), 1% penicillin G/streptomycin (Sigma-Aldrich) at 37 °C under 5% CO₂. Umbilical vein endothelial cells (HUVECs) were maintained in Endothelial cell Basal Medium-2 (EBM-2) (Lonza, Basel, Switzerland) at 37 °C under 5% CO₂.

Protocols and methods. A Nano-LC/MALDI-TOF system was used with three types of IKKβ (wild-type IKKβ, a dominant-positive mutant of IKKβ and a dominant-negative mutant of IKKβ) that were overexpressed in HEK293T cells and immunoprecipitated by a specific antibody. Each sample was reduced with 45 mM of DTT (Wako), alkylated with 100 mM of iodoacetamide (Sigma-Aldrich) and digested with 2000 ng of trypsin (Promega, Madison, WI, USA). One-dimensional peptide fractionation was performed with a Dionex Direct Nano-Flow LC/MALDI system (KYA Tech., Tokyo, Japan) using a reverse-phase (RP) trap column (HiQ SIL C18-3, 0.8 mm i.d. × 3 mm) and an RP analytical column (HiQ SIL C18-3, 0.15 mm i.d. × 30 mm). The samples were subjected to the trap column and then to the analytical column using a gradient of 0–50% solvent B in solvent A over 6 minutes [solvent A: 0.1% trifluoroacetic Acid (TFA), 2% acetonitrile; solvent B: 0.1%TFA, 70% acetonitrile] followed by 50–100% solvent B for 15 minutes at a flow rate 200 nL/minute. The RP column eluent was spotted onto a MALDI sample plate using a Dionex Direct Nano-Flow LC/MALDI system (KYA Tech.) and analyzed using a 4800 mass spectrometer (Applied Biosystems Inc.). Peptides were fragmented under collision-induced dissociation conditions to give fragment ions that produced sequence information for the peptide. The software packages used for data acquisition and analysis included GPS explorer (Applied Biosystems Inc.) and Mascot (Matrix Science, Boston, MA, USA), respectively. The parameters of tolerance for the searches were set to 100 ppm for the MS and 0.2 Da for the MS/MS analyses.

Co-immunoprecipitation (Co-IP). HEK293T cells (8 × 10⁶) were co-transfected with 0.5 µg of AMAP1 plasmids and 0.5 µg of IKKβ plasmids using Effectene reagent (Qiagen, Valencia, CA, USA) according to the manufacturer’s protocol. After 48 h, the cells were washed with PBS and lysed with Cell Lysis Buffer (CST). One-milligram of the lysate was incubated with 1 µg of control IgG, anti-HA tag or anti-FLAG antibody overnight at 4 °C with agitation. Twenty-five microtiter of protein G Sepharose (GE Healthcare, Uppsala, Sweden) was added, and the samples were rotated at 4 °C for 1 h. The beads were washed three times with Cell Lysis Buffer and one time with Tris buffer pH 7.5. Proteins that were bound to the beads were eluted into SDS sample buffer, and the eluted material was analyzed by immunoblotting using anti-HA, anti-FLAG and anti-GST antibodies. Co-precipitation of endogenous AMAP1 with IKKβ was performed using anti-AMAP1 and anti-IKKβ antibodies.

Measurement of p65 activity. HEK293T cells were lysed with NE-PER Nuclear and Cytoplasmic Extraction Reagents (Thermo Scientific) for nuclear and cytoplasmic extract. The transcription factor kits were used to detect the active NF-κB according to the manufacturer’s protocol. Briefly, the biotinylated NF-κB consensus sequence was bound to streptavidin-coated ELISA wells. Because only the active form of NF-κB (p65) binds to the DNA sequence, nonspecific binding was minimized. The p65 that was bound to the consensus sequence was incubated with anti-p65 antibody and then with a secondary, HRP-conjugated antibody. A chemiluminescent substrate was added to the wells, and the resulting signal was detected using a luminometer.

Sucrose gradient. Cells were treated with 5 ng/mL IL-1β or PBS for 30 min and lysed in cell lysis buffer (CST) containing 1 mM PMSF. An identical amount of protein from each sample was mixed with 90% sucrose in MES buffer (50 mM MES, 150 mM NaCl, 1% sucrose concentration 57.5–58.7%) and transferred to a Beckman ultracentrifuge tube. Four milliliters of 35% sucrose followed by 3 mL of 5% sucrose were overlaid, the samples were spun in a Beckman Coulter ultracentrifuge (39,000 rpm; approximately 180,000 × g in a SW40Ti rotor, 20 h), and 26 fractions were collected from the top of the gradient. For the detection of the lipid raft fractions, all fractions were dot-blotted with HRP-labeled cholera toxin B (Sigma-Aldrich) to detect Ganglioside GM1.

Immunofluorescence. HUVECs were grown on a 4-well chamber slide (Thermo Scientific) and treated with or without IL-1β 2.5 mg/mL for 30 min. The cells were washed with PBS, fixed with 4% paraformaldehyde and permeabilized with 0.5% Triton X-100. The cells were blocked incubated with blocking solution (5% goat normal serum in PBS) and labeled with anti-AMAP1 and anti-vimentin antibodies that were coupled to Alexa Fluor 488- and 584-conjugated secondary antibodies. The stained cells were mounted using Mounting Medium with DAPI (Vector Laboratories Inc., Burlingame, CA, USA). Images were captured and exported using a confocal microscope (Olympus FV1000; 60× oil-immersion lens, FLUOVIEW 3.0 software).

Statistical analysis. The data are presented as the mean of ±SD. Groups were compared using a two-tailed, Student’s t-test. P < 0.05 was considered significant.

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Additional information
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