The DUSP26 phosphatase activator adenylate kinase 2 regulates FADD phosphorylation and cell growth

Hyunjoo Kim1*, Ho-June Lee2*, Yumin Oh1, Seon-Guk Choi1, Se-Hoon Hong1, Hyo-Jin Kim2, Song-Yi Lee1, Ji-Woo Choi3, Deog Su Hwang1, Key-Sun Kim3, Hyo-Joon Kim4, Jianke Zhang5, Hyun-Jo Youn6, Dong-Young Noh7 & Yong-Keun Jung1

Adenylate kinase 2 (AK2), which balances adenine nucleotide pool, is a multi-functional protein. Here we show that AK2 negatively regulates tumour cell growth. AK2 forms a complex with dual-specificity phosphatase 26 (DUSP26) phosphatase and stimulates DUSP26 activity independently of its AK activity. AK2/DUSP26 phosphatase protein complex dephosphorylates fas-associated protein with death domain (FADD) and regulates cell growth. AK2 deficiency enhances cell proliferation and induces tumour formation in a xenograft assay. This anti-growth function of AK2 is associated with its DUSP26-stimulating activity. Downregulation of AK2 is frequently found in tumour cells and human cancer tissues showing high levels of phospho-FADDSer194. Moreover, reconstitution of AK2 in AK2-deficient tumour cells retards both cell proliferation and tumourigenesis. Consistent with this, AK2+/- mouse embryo fibroblasts exhibit enhanced cell proliferation with a significant alteration in phospho-FADDser191. These results suggest that AK2 is an associated activator of DUSP26 and suppresses cell proliferation by FADD dephosphorylation, postulating AK2 as a negative regulator of tumour growth.
A denylate kinase (AK) is a ubiquitous enzyme that catalyses the nucleotide phosphoryl exchange reaction. Various isozymes of AK have so far been characterized and are located in different tissues and subcellular region. Among them, AK2 (AK2/ATP–AMP transphosphorylase) is a metabolic enzyme often linked to mitochondrial metabolic rate and has been proposed to have a role in mitochondria–nucleus energetic signal communication. Recent studies have highlighted new exciting developments regarding multi-faceted AK2. AK2 is released into the cytosol to form an apoptotic complex upon initiation of cell death. Moreover, human AK2 deficiency causes a hematopoietic defect associated with sensorineural deafness and Kieloid disease associated with fibroproliferative tumours developing as a result of deregulated wound healing. AK2 was also proposed to associate with mitotic spindle apparently to power cell division centrodes and its dysregulation was observed in human cancers and amyotrophic lateral sclerosis (ALS) diseases. Thus, AK2 may have important extra-mitochondrial functions, especially in tumourigenesis.

Results

Nuclear AK2 negatively regulates FADD phosphorylation. Based on our earlier report showing that AK2 binds to FADD, we addressed whether AK2 is responsible for the regulation of FADD phosphorylation. Reduction of AK2 expression apparently enhanced the level of p-FADD, whereas ectopic expression of AK2 reduced it (Fig. 1a). As reported, we confirmed the increase of this FADD phosphorylation after treating cells with phosphatase inhibitors, okadaic acid and calyculin A, and no phosphorylation of an FADD S194A mutant in which Ser194 was replaced with an Ala (Supplementary Fig. 1a). In addition, western blot analysis following two-dimensional PAGE confirmed that the appearance of only one p-FADD that migrated more slowly to a more acidic pH than non-phosphorylated FADD was exclusively regulated by AK2 (Supplementary Fig. 1b).

Unlike AK2, ectopic expression of cytosolic AK1 or mitochondrial AK3 had no effect on FADD phosphorylation (Fig. 1b). Moreover, a nucleotide kinase-dead mutant, AK2 K28E (ref. 5), reduced FADD phosphorylation as effectively as wild-type AK2 (Fig. 1c). The ability of AK2 to regulate FADD phosphorylation was also observed in Chang liver and Huh-7 tumour cells (Supplementary Fig. 1c). It thus appears that the activity responsible for FADD dephosphorylation is a unique feature of AK2 among AK isotypes and differs from its AK activity.

To better understand a molecular function of AK2, we used a set of AK2 deletion mutants and determined AK2 domain responsible for the regulation of FADD dephosphorylation (Fig. 1d). All of the AK2 constructs were fused with green fluorescence protein (GFP), enabling us to visualize their subcellular localization. From ectopic expression analysis, we found that the AK2 N3 mutant comprising the N-terminal NMPbind (amino-acid residues 45–74) and middle LID (amino-acid residues 141–178) regions exhibited the phosphatase activity against p-FADD, which was comparable to that of wild-type AK2 (Fig. 1d). As seen with the AK2 N2 and AK2 C2 mutants, however, further deletion of the NMPbind domain (AK2 C2) or LID domain (AK2 N2) from the AK2 N3 abolished the enzymatic activity to regulate FADD phosphorylation (Fig. 1d).

Despite its proposed localization in mitochondria, an amino-acid homology search suggests a high probability of nuclear AK2 localization (34.8%; PSORT II programme). We therefore examined subcellular distribution of AK2 using fractionation analysis. Although the majority of AK2 localized in the mitochondrial fraction, a significant amount of AK2 was also detected in the nuclear fraction of HeLa cells (Supplementary Fig. 2a). When we isolated AK2/FADD complexes from nuclear fraction with immunoprecipitation assay and assessed phosphatase activity of the immunoprecipitates using universal substrate para-nitrophenylphosphate (pNPP), the protein complexes containing AK2 or AK2 N3 mutant exhibited higher phosphatase activity than other AK2 mutants (Fig. 1e). These results suggest that AK2-associated phosphates activity is present in the nuclear fraction. In keeping with this notion, when we enforced AK2 to be located exclusively in the mitochondria by adding additional mitochondria-targeting sequence of cytochrome c oxidase subunit VIII, the mitochondria-targeted AK2 failed to reduce the level of p-FADD in cells (Supplementary Fig. 2b). Besides, p-FADD detected in the nucleus was augmented upon short hairpin RNA (shRNA)-mediated repression of nuclear AK2 (Supplementary Fig. 2c).

Next, we determined whether AK2 itself has a phosphatase activity. Human AK2 protein was expressed in and purified from bacteria, after which enzymatic activity of the purified protein was assessed in vitro. However, the purified AK2 protein exhibited no phosphatase activity towards pNPP (Fig. 1f) and a synthetic phospho-Ser-194 peptide (191GAMGSPMS197) derived from the
Figure 1 | AK2 regulates FADD phosphorylation. (a) AK2-dependent regulation of FADD phosphorylation. HeLa cells were transfected with pEGFP (Control), pAK2-GFP or pAK2-shRNA for 24 h and then left untreated or treated with okadaic acid (OA, 0.5 μM) and calyculin A (CA, 50 nM) for 2 h. Cell extracts were analysed by western blotting with the indicated antibodies. (b) Ectopic expression of AK2, but not AK1 or AK3, dephosphorylates FADD. HeLa cells were transfected with pAK1-HA, pAK2-HA or pAK3-HA for 24 h, after which cell extracts were subjected to western blotting with anti-p-FADD, anti-FADD or anti-HA antibodies. (c) The ability of AK2 to dephosphorylate FADD is independent of its enzyme activity. HEK293T cells were transfected with pcDNA (Control), pAK2-GFP, pAK2 K28E-GFP or pAK2 R150A-GFP for 24 h, and cell extracts were then prepared and subjected to western blotting. (d) The N-terminal and middle regions of AK2, comprised of the NMPbind and LID domains (AK2 N3), mediate FADD dephosphorylation. HeLa cells were transfected with pAK2 or AK2 deletion mutants for 24 h, and cell extracts were subjected to western blotting. NS indicates non-specific signal. (e) The AK2 N3 mutant exhibits phosphatase activity. HEK293T cells were co-transfected with p-FADD-HA and either pAK2 or AK2 deletion mutants for 24 h. After fractionation of cell extracts, nuclear fractions were subjected to immunoprecipitation (IP) using anti-HA antibody, after which the immunoprecipitates were analysed for the phosphatase activity using pNPP. (f) Purified AK2 protein does not exhibit a phosphatase activity. Bacterially purified GST, His-AK2, His-AK3 and calf intestinal phosphatase (CIP) proteins were incubated with pNPP for 15 min and the OD405 nm of the reaction supernatants was measured with spectrophotometer. Purified proteins used in these assays were stained with Coomassie-blue (insert). (g) Purified AK2 protein enhances the phosphatase activity of FADD-containing protein complexes. HEK293T cells were transfected with pcDNA-HA or p-FADD-HA for 24 h, after which cell extracts were subjected to IP assay using anti-HA antibody. Half of the immunocomplex was analysed by western blotting with anti-HA antibody (insert) and the other half was incubated with purified GST, His-AK2 or His-AK3 protein (each 10 μg) for 1 h. Then, the reactions were examined for phosphatase activity using pNPP. Bars represent mean ± s.d. (n = 3).

Figure 1 | AK2 regulates FADD phosphorylation. (a) AK2-dependent regulation of FADD phosphorylation. HeLa cells were transfected with pEGFP (Control), pAK2-GFP or pAK2-shRNA for 24 h and then left untreated or treated with okadaic acid (OA, 0.5 μM) and calyculin A (CA, 50 nM) for 2 h. Cell extracts were analysed by western blotting with the indicated antibodies. (b) Ectopic expression of AK2, but not AK1 or AK3, dephosphorylates FADD. HeLa cells were transfected with pAK1-HA, pAK2-HA or pAK3-HA for 24 h, after which cell extracts were subjected to western blotting with anti-p-FADD, anti-FADD or anti-HA antibodies. (c) The ability of AK2 to dephosphorylate FADD is independent of its enzyme activity. HEK293T cells were transfected with pcDNA (Control), pAK2-GFP, pAK2 K28E-GFP or pAK2 R150A-GFP for 24 h, and cell extracts were then prepared and subjected to western blotting. (d) The N-terminal and middle regions of AK2, comprised of the NMPbind and LID domains (AK2 N3), mediate FADD dephosphorylation. HeLa cells were transfected with pAK2 or AK2 deletion mutants for 24 h, and cell extracts were subjected to western blotting. NS indicates non-specific signal. (e) The AK2 N3 mutant exhibits phosphatase activity. HEK293T cells were co-transfected with p-FADD-HA and either pAK2 or AK2 deletion mutants for 24 h. After fractionation of cell extracts, nuclear fractions were subjected to immunoprecipitation (IP) using anti-HA antibody, after which the immunoprecipitates were analysed for the phosphatase activity using pNPP. (f) Purified AK2 protein does not exhibit a phosphatase activity. Bacterially purified GST, His-AK2, His-AK3 and calf intestinal phosphatase (CIP) proteins were incubated with pNPP for 15 min and the OD405 nm of the reaction supernatants was measured with spectrophotometer. Purified proteins used in these assays were stained with Coomassie-blue (insert). (g) Purified AK2 protein enhances the phosphatase activity of FADD-containing protein complexes. HEK293T cells were transfected with pcDNA-HA or p-FADD-HA for 24 h, after which cell extracts were subjected to IP assay using anti-HA antibody. Half of the immunocomplex was analysed by western blotting with anti-HA antibody (insert) and the other half was incubated with purified GST, His-AK2 or His-AK3 protein (each 10 μg) for 1 h. Then, the reactions were examined for phosphatase activity using pNPP. Bars represent mean ± s.d. (n = 3).

AK2 binds to DUSP26 and promotes its interaction with FADD. Data provided above led us to ask whether AK2 might act as a regulatory protein that could stimulate the phosphatase activity responsible for FADD dephosphorylation. To address this point, we collected 107 cDNAs encoding Ser/Thr phosphatase, including PPM phosphatase, PPP phosphatase, Ser/Thr DUSP and pyrophosphatase, and carried out functional screening assays. Consistent with earlier reports, AK2 frequently showed a migration shift on western blots with the slowly migrating band corresponding to p-FADD (Fig. 2a). Through western blot analysis following co-expression of FADD-HA and each of the 107 Ser/Thr phosphatases, we were able to isolate DUSP26 as a strong candidate phosphatase for FADD. Among DUSPs, ectopic expression of DUSP26 reduced the level of p-FADD in HEK293T cells and increased the level of the fast-migrating FADD (Fig. 2a). Expressions of DUSPs used here were confirmed by their enhanced phosphatase activities in cell extracts (Fig. 2a). In contrast to wild-type DUSP26, expression of phosphatase-dead (C152S) DUSP26 mutant had no effect on the level of p-FADD (Fig. 2b). Conversely, knockdown of DUSP26 expression using shRNA increased the levels of p-FADD (Fig. 2c). Moreover, DUSP26 was also detected in the nuclear fraction of HeLa cells (Supplementary Fig. 4). These results indicate that DUSP26 may be a phosphatase responsible for FADD dephosphorylation in growing cells.

In order to clarify a close connection between AK2 and DUSP26, we attempted other approach to identify the proteins that physically interacted with DUSP26. Among DUSPs, ectopic expression of DUSP26 reduced the level of p-FADD in HEK293T cells and increased the level of the fast-migrating FADD (Fig. 2a). Expressions of DUSPs used here were confirmed by their enhanced phosphatase activities in cell extracts (Fig. 2a). In contrast to wild-type DUSP26, expression of phosphatase-dead (C152S) DUSP26 mutant had no effect on the level of p-FADD (Fig. 2b). Conversely, knockdown of DUSP26 expression using shRNA increased the levels of p-FADD (Fig. 2c). Moreover, DUSP26 was also detected in the nuclear fraction of HeLa cells (Supplementary Fig. 4). These results indicate that DUSP26 may be a phosphatase responsible for FADD dephosphorylation in growing cells.

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containing protein complexes (Fig. 2d). Consistently, purified AK2 protein bound to 35S-methionine-labeled DUSP26 in vitro (Supplementary Fig. 5a) and the interaction between over-expressed AK2 and FADD was readily detected in HEK293T cells (Fig. 2e). Further, we found that endogenous AK2 interacted with DUSP26 in HeLa cells and vice versa (Fig. 2f), and this...
interaction was evident especially in the nuclear-enriched fraction of growing cells (Supplementary Fig. 5b). Given that FADD is phosphorylated in G2/M phase of cell cycle, we increased cell population in G2/M phase by treating cells with nocodazole, a microtubule destabilizer, and examined the complex formation. Interestingly, we found that DUSP26 preferentially bound to AK2 and FADD in nocodazole-treated and cell cycle-arrested cells, but not in cells undergoing etoposide-induced cell death (Supplementary Fig. 5c). All of these observations suggest that AK2 forms a novel protein complex with DUSP26 in cells that may depend on cell cycle progression.

To gain insight into the molecular mechanism by which AK2 modulates DUSP26-mediated FADD dephosphorylation, we examined whether AK2 affected the interaction of DUSP26 with its substrate. Immunoprecipitation assays showed that AK2 overexpression enhanced the interaction between DUSP26 and FADD (Fig. 2g), whereas AK2 knockdown markedly reduced it (Fig. 2h). It is also noteworthy that AK2 knockdown resulted in a concomitant accumulation of p-FADD retained in DUSP26 complexes (Fig. 2h). To determine the structural network of AK2/DUSP26 complex and its substrate FADD, we performed domain mapping analysis for their interactions. Detailed binding assays using deletion mutants indicated that the middle region of AK2 containing NMPbin is required for its binding to DUSP26 (Supplementary Fig. S5d) and the N- or C terminus of AK2 interacted with death domain of FADD, whereas DUSP26 bound to death effector domain of FADD (Supplementary Fig. 5e).

These observations suggest that AK2 may also participate in and strengthen the protein–protein interaction between AK2/DUSP26 and FADD, allowing FADD to be better recruited into AK2/DUSP26 complex.

**AK2 directly enhances the phosphatase activity of DUSP26.** We then focused on the regulation of DUSP26 phosphatase by AK2. Based on aforementioned data, we hypothesized that AK2 might directly regulate the phosphatase activity of DUSP26. To address this question, DUSP26 protein was purified from bacteria and subjected to *in vitro* phosphatase assays using pNPP. Significant amount of phosphatase activity was readily detected in the purified DUSP26 protein itself (Fig. 3a). Interestingly, the phosphatase activity of DUSP26 protein was enhanced 2- to 3-fold by addition of AK2 protein, depending on the amounts of AK2 protein added. By contrast, addition of AK3 or FADD protein had no effect on the phosphatase activity of DUSP26 (Fig. 3a). AK2 K28E AK-dead mutant protein also enhanced the phosphatase activity of DUSP26 as much as wild-type AK2 did (Fig. 3b). Furthermore, *in vitro* enzyme assays exhibited that the purified DUSP26 protein reduced the amount of p-FADD and this reaction was stimulated by addition of AK2 protein in a synergistic manner (Fig. 3c). These results suggest that DUSP26 functions as a FADD phosphatase and AK2 directly activates the phosphatase activity of DUSP26 independently of its AK activity.

![Figure 3](image-url) | **AK2 enhances the phosphatase activity of DUSP26 in vitro and in cells.** (a) Purified AK2 protein enhances the phosphatase activity of DUSP26 in *in vitro*. His-DUSP26 protein was purified from bacteria and incubated for 30 min with His-AK2 or His-AK3 protein (each 3 μg). Phosphatase assays were performed with pNPP. Bars represent mean ± s.d. (n = 3). (b) AK-dead mutant enhances the phosphatase activity of DUSP26. His-DUSP26 protein was incubated with GST, GST-AK2, GST-AK2 K28E or GST-AK2 R150A protein for 30 min and then subjected to phosphatase assay. Bars represent mean ± s.d. (n = 3). (c) AK2 facilitates DUSP26-mediated dephosphorylation of p-FADD. His-FADD protein was phosphorylated by Plk kinase in *in vitro* and then incubated at 37°C for 1 h with GST-AK2, His-DUSP26 alone or both GST-AK2 and His-DUSP26 (each 5 μg) in the presence or absence of okadaic acid (OA) and calyculin A (CA). The reaction mixtures were then subjected to western blotting with anti-AK2, anti-FADD, anti-p-FADD and anti-DUSP26 antibodies. (d) Ectopic expression of AK2 enhances DUSP26-catalysed FADD dephosphorylation in cells. HEK293T cells were co-transfected with combinations of the indicated vectors for 12 h, after which cell extracts were subjected to western blotting. (e) Reduction of AK2 expression decreases the phosphatase activity of DUSP26-containing protein complexes. HeLa/Cont shRNA and HeLa/AK2 shRNA cells were transfected with pHA-DUSP26 for 36 h, after which cell extracts were subjected to immunoprecipitation (IP) assay with anti-HA antibody. Halves of the immunoprecipitates were then probed by western blotting with anti-AK2 and anti-HA antibodies and the other halves were assayed for phosphatase activity. Whole-cell lysates (WCL) were included for the comparison of AK2 and HA-DUSP26 expression. Bars represent mean ± s.d. (n = 3). (f) Downregulation of AK2 expression reduces DUSP26-catalysed FADD dephosphorylation. HEK293T cells were co-transfected with p-FADD-HA, pFlag-DUSP26 and either pSuper (Control) or pAK2-shRNA for 24 h, after which cell extracts were subjected to western blotting.
We then characterized the stimulatory effect of AK2 on DUSP26 activity in cells by examining their substrate, p-FADD. Cellular levels of p-FADD, which were significantly reduced by ectopic expression of either DUSP26 or AK2, was severely restrained by co-expression of DUSP26 and AK2 (Fig. 3d). In addition, we found that the reduction of p-FADD by AK2 expression was hampered by the treatment with NSC-87877, a potent inhibitor of DUSP26 (ref. 43) (Supplementary Fig. 6). What’s more, when we isolated DUSP26 complex with immunoprecipitation assays and compared their enzyme activities, the phosphatase activity of DUSP26 complexes isolated from AK2 knockdown cells was twofold lower than that from control cells (Fig. 3e). Under this condition, DUSP26 complex isolated from control and AK2 knockdown cells contained similar amounts of HA-DUSP26 (Fig. 3e). Furthermore, downregulation of AK2 expression weakened DUSP26-mediated FADD dephosphorylation (Fig. 3f). These observations strongly support the notion that DUSP26 activity is positively regulated by AK2.

**Purified AK2 and DUSP26 form a binary protein complex.** To examine the formation of AK2/DUSP26 protein complexes *in vitro*, we carried out a size-exclusion chromatography analysis using Superdex 200. When analysed separately, purified glutathione S-transferase (GST)-AK2 and DUSP26 proteins were detected with their respective molecular weight (His-DUSP26, 26 kDa; and GST-AK2, 56 kDa) (Fig. 4a). Significant phosphatase activity was detected in fraction 36 that contained DUSP26 (Fig. 4a). When incubated together and analysed, however, DUSP26 and AK2 proteins were all detected in two groups of early fractions, fractions 13–17 (∼400 kDa) and 22–26 (∼100 kDa) (Fig. 4b). Compared with fraction 36 that contained only DUSP26, these two groups showed 2- to 3 times higher phosphatase activity (Fig. 4b). Further, we observed formation of a binary complex containing AK2 and DUSP26 in fractions 22 and 24 (Fig. 4b). These *in vitro* results support our proposal that AK2 forms a protein complex with DUSP26 to enhance the phosphatase activity of DUSP26.

**AK2 regulates FADD phosphorylation against cell growth.** Based on the previous reports showing that FADD phosphorylation has been associated with cell cycle progression, we next evaluated whether AK2 was accountable for FADD-mediated cell cycle alternation. Analysis of HeLa cells at different stages of cell cycle with western blotting revealed that the level of p-FADD was higher at early M phase in AK2 knockdown cells than in control cells (Supplementary Fig. 7a). In support of this, depletion of AK2 in HeLa cells led to significant increase of cell proliferation (Supplementary Fig. 7b). We also assessed the role of DUSP26 in cell cycle alteration and cell growth. Consistently, depletion of DUSP26 in HeLa cells resulted in increase of FADD phosphorylation at early M phase (Supplementary Fig. 7c) and facilitated cell proliferation (Supplementary Fig. 7d). These observations suggest that AK2 and DUSP26 have crucial roles in the regulation of cell proliferation.

To address this perspective of AK2 function in detail, we assessed the pathological significance of AK2 function by examining the connection between AK2 dysregulation and cell proliferation in human tumour cells and cancer tissues. Although AK2 expression was detected in most cancer cells, including SK-Hep1 and HepG2 hepatocellular carcinoma cells, its expression was drastically downregulated in MCF-7 breast cancer cells

**Figure 4 | Purified AK2 protein forms a complex with DUSP26 on gel filtration.** (a) Elusion profile and phosphatase activity of each AK2 or DUSP26 protein on size-exclusion chromatography analysis. Purified GST-AK2 or His-DUSP26 protein (each 10 μg) was separated on a Superdex 200 gel filtration column (0.6-ml fractions). Proteins of each fraction were then separated by SDS-PAGE and analysed by western blotting with anti-AK2 or anti-DUSP26 antibody. Each fraction was also assayed for the phosphatase activity using pNPP. (b) The formation of AK2/DUSP26 binary protein complexes on size-exclusion chromatography and stimulation of DUSP26 phosphatase activity by AK2. Purified GST-AK2 and His-DUSP26 proteins (each 10 μg) were combined together for 1h and then subjected to size-exclusion chromatography as in a. Each fraction was examined by western blotting and phosphatase assays. Protein fractions (number 22, 24 and 28) were subjected to GST pull-down assays and the pulled samples were analysed by western blotting with anti-AK2 and anti-DUSP26 antibodies.
and C33A cervical cancer cells (Fig. 5a). In addition, AK2 expression was also downregulated in other HS-578T, MDA-MB-453, BT-474 and SK-BR-3 breast cancer cells among total eight breast cancer cell lines (Fig. 5b). Interestingly, high levels of p-FADD were detected in MCF-7, MDA-MB-453, BT-474, SK-BR-3 and C33A cells, despite there being no difference in total FADD (Fig. 5a,b). Then, we stably reconstituted MCF-7 and C33A cells with AK2 (MCF-7/AK2 and C33A/AK2 cells, respectively) and examined their FADD phosphorylation status and cell proliferation rates. Compared with MCF-7 and C33A cells, the levels of p-FADD were markedly reduced in MCF-7/AK2 and C33A/AK2 cells (Fig. 5c). Moreover, cell proliferation rates of MCF-7/AK2 and C33A/AK2 cells were significantly suppressed by 40% at 6 day (Fig. 5d). Further, treatment with NSC-87877 exerted a strong inhibitory effect on the proliferation of MCF-7/AK2 cells compared with MCF-7 cells (Supplementary Fig. 8). Collectively, AK2 appears to be a strategic controller that acts to coordinate both cell proliferation and FADD dephosphorylation in these tumour cells.

Next we explored whether the cell proliferation–inhibitory activity of AK2 is associated with FADD. When we compared cell proliferation of FADD<sup>−/−</sup> mouse embryo fibroblasts (MEFs) with FADD<sup>+/+</sup> MEFs, FADD<sup>−/−</sup> MEFs displayed slower proliferation than FADD<sup>+/+</sup> MEFs (Fig. 5e). In the similar context, when we introduced AK2 into FADD<sup>−/−</sup> and FADD<sup>−/−</sup> MEFs, AK2 restrained cell proliferation of FADD<sup>+/+</sup> MEFs but not that of FADD<sup>−/−</sup> MEFs (Fig. 5e). In addition, ectopic expression of AK2 or AK2 N3 mutant, which retained high activity for FADD dephosphorylation, exhibited growth-suppression activity (Supplementary Fig. 9a). Further, we monitored the effect of p-FADD on cellular proliferation using D4476, a selective inhibitor of CK1 kinase, that is responsible for FADD phosphorylation<sup>22</sup>. Treatment with D4476 reduced the level of p-FADD and suppressed cell proliferation in AK2 knockdown cells (Supplementary Fig. 9b), reversing the stimulatory effect of AK2 knockdown on FADD phosphorylation and cell proliferation. As illustrated in Fig. 5f, it thus appears that AK2 exhibits cell proliferation–inhibitory activity in a FADD-dependent manner.

**AK2 suppresses tumourigenesis and is lost in human cancers.**

These observations led us to address whether or not AK2 expression affected tumour growth in vivo. After subcutaneous injection of wild-type cells and AK2 over-expression MCF-7 or C33A cells into the flanks of nude mice, we monitored tumour growth for 12 weeks. Compared with control cells, ectopic restoration of AK2 in MCF-7 and C33A cells lacking AK2 expression led to near-complete tumour regression (Fig. 6a–c). Reversely, ablation of endogenous AK2 expression significantly promoted tumour growth in the right side of mouse flank with 70% of substantial tumour progression (Supplementary Fig. 10a). Analysis of the dissected tumours showed that the inoculation of AK2 knockdown cells into nude mice led to a significant increase of tumourigenesis (Supplementary Fig. 10b–d). These results suggest that downregulation of AK2 expression produces impressive in vivo tumour efficacy in mice.
AK2 expression was reduced in western blot analysis, we confirmed that AK2 expression was decreased in C33A/AK2 and MCF-7/AK2 cells when we expressed AK2 WT or AK2 K28E mutant in MEFs, the phosphorylation of mouse FADD at Ser191 increased (Fig. 7c). These results support that as in human tumour cells, AK2 regulates FADD phosphorylation and cell proliferation in MEFs, showing a conserved role of AK2 in the regulation of cell proliferation and FADD phosphatase activity in mouse tissue.

**Discussion**

Functional screening seeking for AK2-associated phosphatase enabled us to identify DUSP26. Our study on it provides several lines of evidence that AK2 is required as an activator for optimal phosphatase activity of DUSP26. First, downregulation of either AK2 or DUSP26 expression enhances FADD phosphorylation. Second, AK2 directly forms protein complex with DUSP26 and enhances the phosphatase activity of DUSP26. Third, AK2 knockdown reduces the phosphatase activity of DUSP26 in cells. Fourth, tumour cells lacking AK2 exhibit high level of p-FADD and reconstitution of AK2 reverses it. Thus, we propose that AK2 is the first positive regulator of DUSP26 phosphatase. AK2/DUSP26 complex described here, however, forms to regulate cell proliferation and is distinct from AK2/FADD/caspase-10 apoptotic complex in that caspase-10 is not present in AK2/DUSP26 complex. Given that DUSP26 functions as a phosphatase of other substrates, including MAP kinases and p53 (refs 14,15,17,19), it is conceivable that the substrate specificity and activity of DUSP26 are variable by and depend on its associated regulator, such as AK2. In addition, the discrepancy of early reports showing distinct roles of DUSP26 in either stimulation17 or inhibition19–21 is conceivable that the substrate specificity and activity of DUSP26 are variable by and depend on its associated regulator, such as AK2. In addition, the discrepancy of early reports showing distinct roles of DUSP26 in either stimulation17 or inhibition19–21 is consistent with a role of AK2 in cells analysed.

Other functions of AK2 than AK activity regulate the phosphatase activity of DUSP26. Unlike AK1, the enzymatic activity of AK2 is much low5. Then, an important mechanistic question that remains is how AK2 regulates the phosphatase activity of DUSP26. AK2 alone was enough to stimulate DUSP26 activity in vitro. It, thus, appears that the binding of AK2 to DUSP26 may induce conformational change of DUSP26 to
FADD phosphorylation was examined by immunoblotting. (transfected with wild-type AK2-GFP or AK2 K28E-GFP mutant for 24 h and mutant reduces FADD phosphorylation. (and AK2 is also required for DUSP26 to regulate these proteins. to western blotting with anti-mouse pSer191-FADD and anti-AK2 antibodies. AK2 augmented in mitochondria, such as nucleus, regulates cell proliferation. As injection. Therefore, it appears that AK2 present in the non-
localization signals located in its middle (residues 74–141) and also found in the nucleus. We identified at least two nuclear opposite role of AK2 in FADD trafficking. Although AK2 is unclear; however, FADD trafficking requires phosphorylation of the protein on Ser194 (ref. 46). As predicted, AK2 knockdown uncovers; however, FADD trafficking requires phosphorylation of their substrate, FADD. Dysregulation of this AK2/DUSP26 FADD signalling pathway does not have to be limited to breast cancer. We also observed significant downregulation of AK2 in other cancers, such as liver and lung cancers. We, therefore, propose that AK2 is a negative regulator of tumour growth that activates DUSP26 to suppress cell proliferation by FADD in human cancers (Fig. 7d).

Methods

Cell culture and antibodies. HEK293T, C33A and MCF-7 cells (The American Type Culture Collection (ATCC), Manassas, VA, USA) were cultured in Dulbecco’s Modified Eagles Medium (DMEM; HyClone, Logan, UT, USA) supplemented with 10% fetal bovine serum (HyClone). Wild-type and FADD knockout MEFs were cultured with DMEM with 10% fetal bovine serum. Polyclonal rabbit anti-AK2 (Santa Cruz Biotechnology, CA, USA; sc-80070), anti-β-tubulin (Sigma-Aldrich, St Louis, MO, USA), and anti-Flag (Sigma-Aldrich; F8014) antibodies were used in western blot analysis (1 μg ml⁻¹); Transfection was carried out with LipofectAMINE reagent (Invitrogen, Carlsbad, CA, USA) following the manufacturer’s instructions. For generation of stable cell lines, HEK293T cells were transfected with vector using LipofectAMINE reagent (Invitrogen) for 24 h and then grown in selection medium containing 1 mg ml⁻¹ G418 (Invitrogen) for 2 weeks. C33A and MCF-7 cells were also transfected with vector and selected for 2 weeks in the presence of 1.2 mg ml⁻¹ G418. After single cell cloning, the clones were screened by western blot analysis. HEK293T cells were used for the screening and immunoprecipitation assays for maximum efficiency of DNA transfection and the other tumour cells were analysed for cell proliferation.

Plasmid constructs. Construction of pAK2 shRNAs was described previously. For construction of pDUSP26 shRNA, the forward and reverse 60-nucleotide fragments containing sequences of human DUSP26 (shRNA no. 1–5, 5’-gatcgcagaaatgatattgctttcagctatcatgcttcttta-3’, shRNA no. 1–3, 5’-gactttaaaagagacctgacagagagcatctctgttactgcatctgctg-3’, shRNA no. 2–5, 5’-gatccccccatcagaaatgatattgctttcagctatcatgcttcttta-3’, shRNA no. 2–3, 5’-agcttttaaaagagacctgacagagagcatctctgttactgcatctgctg-3’, shRNA no. 3–5, 5’-gatccaccccatcagaaatgatattgctttcagctatcatgcttcttta-3’, shRNA no. 3–2, 5’-agcttttaaaagagacctgacagagagcatctctgttactgcatctgctg-3’, shRNA no. 3–5, 5’-gatccaccccatcagaaatgatattgctttcagctatcatgcttcttta-3’, shRNA no. 4–3, 5’-agcttttaaaagagacctgacagagagcatctctgttactgcatctgctg-3’, shRNA no. 4–2, 5’-agcttttaaaagagacctgacagagagcatctctgttactgcatctgctg-3’) were synthesized, annealed and cloned into the BglII and HindIII sites of pSuper mammalian expression vector (OligoGene, Seattle, WA, USA). The cDNAs encoding AK1, AK2 and AK3 were obtained by PCR and cloned into pcDNA3-HA. The cDNAs encoding AK2 FL, AK2 N1, AK2 N2, AK2 N3, AK2 C1, AK2 C2 and AK2 C3 were cloned into the Xhol site of pEGFP (enhanced green fluorescent protein) N1. DUSP26 cDNA was also obtained by PCR and cloned into the EcoRI and EcoRV sites of p3 × FLAG-CMV-10 vector (Sigma-Aldrich).

Identification of DUSP26-binding proteins by LC-MS/MS. DUSP26-binding proteins were affinity-purified from extracts of HEK293T cells stably expressing 3 × Flag-tagged DUSP26. The DUSP26-binding proteins were immunoprecipitated using anti-Flag antibody-conjugated agarose beads (Sigma-Aldrich) from extracts that were washed with buffer containing 20 mM Tris–HCl, pH 7.9, 15% glycerol, 1 mM EDTA (Fa, 1 mM dithiothreitol (DTT), 0.2 mM phenylmethylsulfonyl fluoride (PMSF), 0.05% Nonidet P40 and 150 mM KCl to remove non-specific contaminants. The bound proteins were eluted by competition with the Flag peptide (0.1 mg ml⁻¹), resolved by SDS–PAGE and prepared for LC-MS/MS analysis. Peptide samples were injected into a column by a Surveyor autosampler (Surveyor; Thermo Finnigan, San Jose, CA, USA) and separated by C18 column. The eluent was directly transferred to the electrospray ionization source of a Thermo Finnigan LCQ DecaXPlus ion trap mass spectrometer. Automated peak recognition,
dynamic exclusion, and daughter ion scanning of the two most intense ions were performed and analysed by the XCALIBUR software.

**In vitro FADD phosphatase assay.** His-FADD was purified from bacteria and incubated with PK in a kinase reaction buffer (20 mM Tris–HCl, pH 7.5, 10 mM MgCl₂, 0.01% Triton X-100, 0.5 mM Na₃VO₄, 2.5 mM DTT and 200 nM ATP). The reaction products were then incubated with purified GST-AK2 or His-DUSP26 (5µg) in the phosphatase reaction buffer (50 mM Tris–HCl, pH 7.4, 150 mM NaCl, 5 mM MgCl₂ and 0.02% Triton X-100). After 1 h at 37°C, the reaction products were separated by SDS-PAGE and analysed by western blot analysis.

pNPP hydrolysis assay. Samples were incubated with pNPP (Sigma-Aldrich) in reaction buffer (25 mM HEPEs, pH 7.2, 50 µM DTT and 2.5 mM EDTA) for 30 min at 25°C and then centrifuged briefly. The pNPP hydrolysis was assessed by measuring the OD₄₀₅nm of the reaction supernatant with spectrophotometer.

**Cytosolic-nuclear fractionation assay.** For separation of the cytosolic or nuclear fraction, HeLa and HEK293T cells were harvested, resuspended in buffer A (25 mM Tris–HCl, pH 8.0, 10 mM KCl, 1 mM DTT and 0.5 mM PMSF) and incubated for 15 min on ice. After cell lysis elicited by the addition of 0.5% NP-40 and vigorous vortexing, the lysates were spun down at 1,400 g for 1 min, and the supernatants were saved as the cytosolic fractions. Nuclear fractions were obtained by resuspending and vortexing the pellets in buffer C (50 mM Tris–HCl, pH 8.0, 400 mM NaCl, 1 mM DTT and 1 mM PMSF) followed by incubation for 30 min on ice. After centrifugation at 15,000 g for 30 min at 4°C, the supernatants were collected as the nuclear fractions.

**RT-PCR.** Total cellular RNA was isolated from cells using Trizol reagent (MRC, Cincinnati, OH, USA). The following primer sets for DUSP26 were then used for in vitro amplification of the mRNA: forward primer 5′-gaaggtgaaggtcggagtca-3′ and reverse primer 5′-gcctacacaggctaa-3′.

**Chromatographic methods.** HeLa cells were lysed by homogenization in hypertonic buffer (50 mM PIPES, 2 mM EDTA, 5 mM DTT, 15 mM MgCl₂, 50 mM KCl, 0.1% CHAPS and protease inhibitors). Cells extracts were then fractionated by size-exclusion chromatography on Superdex 200 analytical columns using a fast protein liquid chromatography protein purification system (Amersham Pharmacia Biotech, Herts, UK). Superdex columns were eluted at 4°C with buffer C (50 mM Tris–HCl, pH 7.0) containing 5% (v/v) sucrose, 0.1% (v/v) CHAPS, 20 mM HEPEs/NaOH and 5 mM DTT. The columns were calibrated with protein standards (Amersham Pharmacia Biotech).

**Generation of AK2 knockout mice.** AK2 gene-trapped embryonic stem cells (RRI133) were provided by BayGenomics of the International Gene Trap Consortium. AK2 heterozygous mice was generated by following the protocol provided by BayGenomics. Genotypes were analysed using genomic PCR with primers provided by BayGenomics. Genotypes were analysed using genomic PCR with genomic DNA from tail biopsies.

**Cell growth assay.** To measure cell density using DNA content, 3×10⁵ cells were seeded into 96-well plates, after which cells were removed every day and immediately frozen at −70°C. Cells were then thawed and lysed, and DNA content was measured using a CyQUANT Cell Proliferation Assay Kit (Invitrogen) by measuring the fluorescent signal (excitation, 480 nm; and emission, 520 nm).

**In vitro-binding assay.** Purified GST-fusion proteins coupled to Glutathione-Sepharose 4B (Amersham Biosciences) were incubated for 3 h in binding buffer (50 mM Tris–HCl (pH 8.6), 100 mM NaCl, 1 mM DTT, 2 mM EDTA, 0.05% Triton X-100, 1 mM PMSF, and 5% (v/v) glycerol) at 4°C with [3H]-methionine-labeled proteins which were in vitro translated using a TNT-coupled transcription/translation system (Promega, Madison, WI, USA). Typically, 20 µg of the immobilized fusion protein was incubated with 10 µl of in vitro translated sample. After pull-down assay with Glutathione-Sepharose beads, the precipitated proteins were extensively washed with binding buffer, separated by 12% SDS–PAGE, and detected by autoradiography.

**Two-dimensional gel electrophoresis.** Cells were washed three times with ice-cold PBS, after which total cell lysates were prepared by sonication in ice-cold buffer (9 M urea, 2% CHAPS, 0.8% pharmalyte and 1% DTT). The lysates were centrifuged at 15,000 g for 30 min, after which the supernatant was collected and stored at −80°C. Samples were applied to 18-cm long pH 4–7 IPG strips (GE Healthcare, Piscataway, NJ) for 12 h at 10 V using IPGphor (GE Healthcare). Isoelectric focusing was performed for 1 h at 500 V and for 1 h at 1,000 V, after which the voltage was gradually increased to 8,000 V for 1 h. After focusing at 8,000 V for an additional 4 h, each strip was incubated for 20 min in buffer containing 50 mM Tris–HCl (pH 8.8), 6 M urea, 30% glycerol, 2% SDS and 1% DTT, and then for an additional 4 h in the same buffer containing 2.5% iodoacetamide (IAA) instead of DTT. The equilibrated strip was placed onto 10–15 gradient polyacrylamide gel and the gel was electrophoresed at 70 V in running buffer (25 mM Tris–HCl, pH 8.8, 192 mM glycine and 0.1% SDS).

**In vitro phosphatase assay using HPLC.** Synthetic FADD (GAMPSMS) and p-FADD peptides (GAMPSMS) (each 20 µg) (Peptron Inc., Daejeon, Korea) were used as substrates for purified AK2 protein (8 µg) for 30 min at 37°C. Then, the reaction products were analysed by HPLC reverse phase.

**FACS analysis.** HeLa cells were spun down, washed with PBS and fixed in 70% ethanol. After incubation for 10 min on ice, cells were centrifuged at 800 g for 10 min at 4°C and then washed three times with ice-cold PBS. The pellets were resuspended in 500 µl of PBS and stained with 50 µg ml⁻¹ propidium iodide (Sigma-Aldrich) in the presence of 100 µg ml⁻¹ RNase (Invitrogen). Samples were then analysed using a FACS calibur flow cytometer (BD Biosciences).

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