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Loss of the HPV-Infection Resistance EVER2 Protein Impairs NF-κB Signaling Pathways in Keratinocytes

Françoise Vuillier1*, Guillaume Gaud1, Delphine Guillemot1, Pierre-Henri Commere2, Christian Pons1, Michel Favre1

1 Unité de Génétique, Papillomavirus et Cancer Humain, Institut Pasteur, Paris, France, 2 Plateforme de cytométrie en flux, Institut Pasteur, Paris, France

Abstract

Homozous mutations in EVER genes cause epidermodysplasia verruciformis (EV), characterized by an immune defect and the development of skin cancers associated with β-human papillomavirus (HPV) infections. The effects of EVER protein loss on the keratinocyte immune response remain unknown. We show here that EVER2 plays a critical role in the interplay between the NF-κB and JNK/AP-1 signaling pathways. EVER2-deficient cells overproduce IL-6 following the upregulation of JNK activation. They respond poorly to phorbol ester and TNF via the NF-κB pathway. They have lower levels of IKKα subunit, potentially accounting for impairments of p100 processing and the alternative NF-κB pathway. The loss of EVER2 is associated with an unusual TRAF protein profile. We demonstrate that EVER2 deficiency sustains TRAF2 ubiquitination and decreases the pool of TRAF2 available in the detergent-soluble fraction of the cell. Finally, we demonstrate that EVER2 loss induces constitutive PKCθ-dependent c-jun phosphorylation and facilitates activation of the HPV5 long control region through a JNK-dependent pathway. These findings indicate that defects of the EVER2 gene may create an environment conducive to HPV replication and the persistence of lesions with the potential to develop into skin cancer.

Introduction

Cutaneous human papillomavirus (HPV) can induce diverse skin lesions, from warts to fully invasive carcinomas [1,2]. The host genetic factors favouring the malignant transformation of oncogenic HPV-infected keratinocytes have not been fully elucidated. The first reported evidence for this association was obtained from patients with epidermodysplasia verruciformis (EV) [3]. This rare autosomal recessive skin disease (OMIM 226400) is associated with abnormally high susceptibility to β-HPVs. EV patients have disseminated skin lesions and frequently develop squamous cell carcinoma induced by HPV5 and 8 [1,4]. They have defective cell-mediated immunity, resulting in the persistence of lesions and high loads of the infecting β-HPVs [2]. EV thus constitutes a model of genetic skin cancer induced by specific HPVs [3].

We have demonstrated that homozygous mutations of EVER1 (TMC8) and EVER2 (TMC6) are associated with EV [5]. These genes are expressed in hematopoietic and endothelial cells [6], consistent with a role in immunity to HPV [2]. The EVER proteins are probably involved in signal transduction pathways in keratinocytes, but their exact role remains unclear. They may regulate zinc homeostasis and the activity of the transcription factor AP-1, a key element in the HPV life cycle [7].

Many studies have provided evidence of a link between NF-κB activation and hematological and epithelial cancers [8,9]. There is also evidence to suggest that NF-κB inhibition increases the incidence of liver and skin cancer, implying cell-specific effects [9,10,11]. The over-production of TNF and TGF-β in the lesional epidermis of EV patients [4] suggested a possible role for inflammatory responses in regulating the growth and differentiation of HPV-infected keratinocytes and in lesion persistence. EVER proteins may thus be involved in regulating the NF-κB and JNK/AP1 pathways and may contribute to HPV-induced carcinogenesis in keratinocytes. We investigated the molecular processes underlying the cancer progression associated with β-HPV infection in the genetic disorder of EV, by assessing the impact of EVER2 loss on the NF-κB and JNK activation pathways, using wild-type and EVER2-deficient keratinocytes. We found that EVER2 loss induced constitutive JNK activation, promoting HPV5 LCR activation and inflammatory responses and highlighting the crucial role of EVER2 in control of the NF-κB and JNK/AP1 signaling pathways.

Results and Discussion

EVER2 deficiency is associated with an abnormal pattern of IL-6 production

We investigated the possible functional consequences of the homozygous truncating mutations of EVER2 causing EV [5], in keratinocyte cell lines lacking EVER2. We generated EVER2−/− keratinocyte cell lines from an EV patient (EV cell line), and from a healthy subject, by silencing EVER2 expression with microRNA (miEVER2 cell line). Wild-type EVER2 cell lines were generated from a healthy subject and are referred to as the “Healthy cell...
EVER2 deficiency impairs NF-κB signaling pathways

TNF and PMA-ionomycin use different upstream pathways converging on IKK activation to stimulate NF-κB [14] (Fig. S1). The changes in luciferase activity in response to these two stimuli led us to assess phosphorylation of the IKK complex upon stimulation. IKK2β complex phosphorylation levels were lower in EVER2−/− cells than in controls following stimulation with PMA-ionomycin (Fig. 3A and S2A). Furthermore, IKKβ phosphorylation was found to be defective following TNF stimulation in EVER2−/− cells (Fig. 3B and S2B). These results indicate that EVER2 deficiency impairs NF-κB signaling pathways.

The abnormal production of IKKζ and IKKβ is associated with skin inflammation, and poor IKKζ production has been reported to promote skin tumor development [15,16,17,18]. We therefore investigated whether EVER2 controlled the levels of IKKζ and β subunits, by assessing the amounts of IKKζ and β in our cell lines. Unlike IKKβ, IKKζ was less abundant in EVER2−/− cells than in control cells, even after PMA-ionomycin stimulation (Fig. 3A and S2A). Thus, EVER2 deficiency impairs the IKKβ complex by decreasing the amount of IKKζ.

Previous studies have suggested that the most important function of IKKζ is the control of p100 processing, leading to activation of the alternative pathway [19] (Fig. S1). We therefore investigated the effects of EVER2 knockdown on the processing of p100 to p52. Constitutively produced p52 was less abundant in the cytoplasm and nucleus of EVER2−/− cells than in those of control cells (Fig. 3C and S2C). After exposure to TNF, p52 levels remained lower in both compartments of EVER2−/− cells than in those of control cells. A defect in basal p52 production was confirmed in the nucleus of healthy cells treated with siRNAs targeting exons 5, 6, 8 and 10 of the EVER2 gene (Fig. S3B). Consistent with the low levels of IKKζ, these findings indicate that EVER2 loss prevents p100 processing, thereby impairing the alternative NF-κB signaling pathway.

The IKKβ subunit is involved principally in activation of the classical NF-κB pathway and it can replace IKKζ if this subunit is missing [16]. We then investigated the effect of EVER2 deficiency on the classical NF-κB signaling axis, by assessing IkBζ levels before and after exposure to PMA-ionomycin or TNF (Fig. S1). No IkBζ degradation was detected in EVER2−/− cells in the basal state, and that following both stimulations was less marked than in control cells (Fig. 3A, 3B, S2A and S2B). As p65 translocation may occur without IkBζ degradation [20], we assessed the p65 subunit in the cytoplasm and nuclear fractions of all cell lines by western blotting before and after stimulation with TNF only (Fig. S1, 3B and S2B), and assessed the p65 translocation by determining the nuclear to cytoplasmic ratio (as illustrated for miEVER2 cell line; Fig. 3D). The amounts of p65 and its phosphorylated form (pH65) translocated into the nucleus were smaller in miEVER2 cells than in controls (Fig. 3D). This translocation defect was confirmed by quantifying p65 and pH65 in nuclear extracts from miEVER2 cells and controls (Fig. 3E). Nevertheless, the translocation defect occurred shortly after stimulation (from 10 min onwards) (Fig. 3D). After stimulation with TNF for 1 hour, the amounts of p65 and its phosphorylated form in the nucleus were significantly lower in EVER2−/− cells than in control cells (Fig. 3E). Immunofluorescence experiments confirmed a translocation defect of p65 into the nucleus in EVER2−/− cells following TNF stimulation. We observed a diffuse staining in cytoplasmic and nuclear compartments of EVER2−/− cells as opposed to control cells where an intense labelling of nuclei was observed (Fig. 31 and S2D). Fluorescence microscopy analysis confirmed also that the p65 subunit was less...
Figure 1. Characterization of cell lines. (A) EV and Healthy cell lines were compared with fibroblasts. They were studied by phase-contrast microscopy. Bars, 50 μm. They were also stained for cytokeratin (KL1); the nucleus was stained with DAPI. Bars, 100 μm. (B–C) EVER2 expression was...
determined in EVER2−/− cell lines (EV and miEVER2) compared with control cell lines (Healthy and miCTRL). EVER2 transcripts (B) were studied by semiquantitative RT-PCR. Healthy cells transfected with control siRNA (siCTRL) were also compared with cells transfected with siRNAs targeting various exons (5, 6, 8 and 10) of EVER2 (siEVER2). The levels of transcripts after various numbers of cycles are shown for RT-PCR studies. GAPDH transcripts were used as the reference. The production of EVER2 protein (C) was evaluated by the western blotting of insoluble fractions isolated from the cells. The data shown are representative of three independent experiments. (D–E) IL-6 production was studied in EVER2−/− (EV and miEVER2) and wild-type (Healthy or miCTRL) cells. The cells were cultured in the presence or absence of TNF. (D) After 48 h, the supernatants were collected and tested for IL-6 and IL-8 by flow cytometry. Data are means ± SD of three independent experiments. (E) After 16 h in the presence of brefeldin A, whole-cell lysates were assayed for intracellular IL-6 by western blotting. Data are from one experiment representative of three independent experiments carried out. *, P<0.05; **, P<0.01; ***, P<0.001.

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Figure 2. EVER2 regulates the transcriptional activity of NF-κB. (A–B) Keratinocyte cell lines were transiently transfected with 0.3 μg/well NF-κB luciferase reporter. The cells were serum-starved overnight and left untreated or were stimulated with TNF (A) or a combination of PMA+ionomycin (B). The cells were harvested and assayed for luciferase activity. The data shown are expressed as fold-induction with respect to nonstimulated cells. Values obtained with wild type cells were taken as 100%. The data shown are means ± SD of three independent experiments. (C) EV or healthy cells were transiently transfected with the NF-κB/luc construct (0.3 μg/well) and various amounts of a plasmid encoding EVER2 or an empty vector as a control. Cells were assayed for luciferase activity after stimulation with TNF following serum starvation. The data shown are expressed as fold-induction with respect to nonstimulated cells. Values obtained with healthy cells transfected with empty vector were taken as 100%. The data shown are means ± SD of three independent experiments. *, P<0.05; **, P<0.01; ***, P<0.001.

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Figure 3. Function of EVER2 in NF-κB and c-jun activation. miEVER2 and miCTRL cell lines were left untreated or were stimulated with PMA+ionomycin (A) or TNF (B–J) for the times indicated. (A) Whole-cell lysates were analyzed by western blotting to explore the IKKα/β complex. The
phosphorylated forms of IKKα and IKKβ were indicated as phIKKα, phIKKβ, respectively. Whole-cell lysates were analyzed for p65 and c-jun using Western blot analysis. The phosphorylated forms of p65 and c-jun are indicated as p65p and c-jun respectively. The phosphorylation of p65 and c-jun was quantified by densitometry. p65 and c-jun levels were set to 1. The results for p65 and c-jun were expressed as the ratio of nuclear and cytoplasmic concentrations. The results for basal levels were set to 1. Data were expressed as the means ± SD of three independent experiments. **P < 0.001, ***P < 0.001.

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**EVER2** deficiency impairs TRAF protein production and sustains TRAF2 ubiquitination

**EVER2** deficiency impairs TRAF protein production and sustains TRAF2 ubiquitination. Western blotting revealed no abnormality in cytoplasmic extracts from **EV** cells than in control cells (Fig. 4D). Immunohistochemistry experiments confirmed the abnormal localization and abundance of TRAF2 in EV cells compared with controls (Fig. 4E). The presence of larger amounts of TRAF2 in nuclear extracts was also confirmed in normal cells after the silencing of **EVER2** with siRNAs specifically targeting exons 5 and 10 (Fig. S3B). We then investigated whether **EVER2** loss affected the level of soluble TRAF2 required for the subsequent stimulation by TNF by assessing the amounts of TRAF2 in Triton-soluble and -insoluble fractions of cells at steady state. EV cells had larger amounts of TRAF2 in insoluble fractions than control cells (Fig. 4F). This abnormal localization was associated with a reduction of TRAF2 in detergent-soluble fraction. In addition, in the insoluble fraction of EV cells, TRAF2 appeared to be a doublet, with one form migrating more rapidly. This may ultimately lead to degradation of TRAF2. Thus, **EVER2** loss induced an accumulation of TRAF2 in detergent-insoluble fraction and reduced the pool of TRAF2 available in the soluble fraction. We therefore investigated whether **EVER2** could interact with TRAF1 or TRAF2, regulating the functions of these proteins. Co-immunoprecipitation experiments showed that **EVER2** formed a complex with both TRAF1 and TRAF2 (Fig. 4G). By contrast, no complex was detected with the receptor-interacting serine/threonine-protein kinase 1 (RIPK1) confirming the specificity of the binding to **EVER2** (Fig. 4G).

The polyubiquitination of TRAF2, a downstream target of PKC, has been shown to contribute to IKK and JNK activation [24]. We therefore investigated whether **EVER2** affected TRAF2 ubiquitination. TRAF2 was constitutively ubiquitinated in **miEVER** but not in control cells (Fig. 4H). For confirmation of the effect of **EVER2** on TRAF2 ubiquitination, we performed TRAF2-immunoprecipitation assays in HEK-293T cells following the overproduction of **EVER2** together with TRAF2 and ubiquitin. The overproduction of **EVER2** and TRAF2 resulted in much lower levels of the ubiquitinated forms of TRAF2 following TNF stimulation (Fig. 4I). Thus, **EVER2** loss sustains constitutive TRAF2 ubiquitination consistent with basal JNK
Figure 4. EVER2 deficiency impairs TRAF protein production. (A–D) EV and Healthy cell lines were left untreated or stimulated with TNF for the time periods indicated. (A) Cytoplasmic extracts were subjected to western blotting. The results shown are representative of three independent experiments. (B) Whole-cell lysates were analyzed by western blotting after various times. (C) qRT-PCR was used to measure the induction of TRAF1 1.5–12 h after stimulation with TNF. Results are reported as fold-induction with respect to untreated cells. The basal level for the Healthy cell line was
set to 1. A logarithmic (log) scale was used for the Y axis. Experiments were performed twice independently. (D) Nuclear extracts were subjected to immunoblot analysis with TRAF2 antibody for the times indicated. The results shown are representative of three independent experiments. (E–F) EV and Healthy (Heal.) cell lines were left untreated. (E) Cells were stained for TRAF2; the nucleus was stained with DAPI. Bars, 20 μm. (F) Keratinocytes were lysed in Triton X-100 lysis buffer. Soluble and insoluble fractions were subjected to immunoblot analysis with TRAF2 antibody. The TRAF2 and actin bands were quantified by densitometry for the soluble fraction. Results are reported as the ratio of TRAF2 to actin. The ratio obtained for the Healthy cell line was set to 1. (G) HEK-293T cells were transfected with plasmids encoding Flag-tagged EVER2 and Gluc2-tagged TRAF1, TRAF2 or RIPK1 proteins or Gluc2 fragment as a control (Gluc2-Ø). Crude lysates were immunoprecipitated with anti-Flag antibody (IP) and immunoblotted (IB) with the antibodies indicated. (H) Endogenous TRAF2 was immunoprecipitated (IP) with anti-TRAF2 in miEVER2 and control cells, and then subjected to immunoblotting (IB) with anti-Ubiquitin (Ub) antibody. The results shown are representative of three independent experiments. (I) HEK-293T cells were cotransfected with Gfp-tagged TRAF2, hemaglutinin (HA)-tagged Ubiquitin (Ub) with or without Flag-EVER2. After 24 h, the cells were treated with TNF for 2 h. Lysates were subjected to immunoprecipitation and immunoblotting with the indicated antibodies. The results shown are representative of two independent experiments.

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Figure 5. EVER2 controls NF-κB and JNK/AP-1 pathways. (A–D) Cell lines were left untreated (-) or were treated with specific inhibitors of JNK (SP60025 (SP; 20 μM)), PI3K (Ly294002 (Ly; 15 μM)), IκBα (Bay11-7082 (Bay; 5 μM)), PKCα (Go6976 (Go; 0.5 μM)), AKT1/2 (Akti; 2 μM), EGF receptor (AG1478 (AG; 12.5 μM)) or N-acetylcysteine (NAC; 10 μM) as indicated. EV and Healthy cell lines were left unstimulated (A) or were stimulated with TNF (B) in the presence of brefeldin A for 16 hours. Whole-cell lysates were assayed for intracellular IL-6 by western blotting. The data shown are from one experiment representative of two independent experiments carried out. The IL-6 and actin bands on western blots were quantified by densitometry. Results are reported as the ratio of IL-6 to actin. In panel A, the ratio obtained for untreated EV cells was set to 1. In panel B, the ratio was set to 1 for each cell line in the absence of inhibitors. (C) miEVER2 and miCTRL cell lines were left untreated (+) or were treated with inhibitors specific as indicated. We analyzed the levels of c-jun or its phosphorylated form (phc-jun) in nuclear extracts by western blotting. The results shown are representative of two independent experiments. (D) EV and Healthy cell lines were transiently transfected with 0.3 μg/well HPV5-flLCR/luc or HPV5-minLCR/luc. Cells were assayed for luciferase activity. Results are reported as fold-increases over the level obtained with HPV5-minLCR/luc. Values obtained with the unstimulated Healthy cell line are set to 100%. The data shown are means ± SD of three independent experiments. *, P<0.05; **, P<0.01; ***, P<0.001. doi:10.1371/journal.pone.0089479.g005

EVER2 deficiency drives an inflammatory response by promoting JNK activation and sustains the PKCα-dependent phosphorylation of c-jun

We observed an unusual amount of IKKα at the basal level (Fig. 3A time 0 and Fig. S2A time 0) associated with a defective response to TNF through NF-κB pathways in EVER2−/− cells (Fig. 3 panels B to G, I and J and S2 panels B to E). We thus carried out inhibition assays, to elucidate the signaling pathways involved in IL-6 production in EVER2-deficient cells (Fig. 1E).
Constitutive and TNF-induced IL-6 production in EV cells was decreased by SP60025 and, to a lesser extent, by Bay11-7082 and Ly294002, specific inhibitors of JNK, IKKβ, and PI3K, respectively (Fig. 5A and 5B top panel). By contrast, no basal IL-6 production was detectable in the Healthy cell line (Fig. 1E and 5A), in which TNF-induced IL-6 production was largely inhibited by treatment with Bay11-7082 and to a lesser extent by Ly294002 (Fig. 5B bottom panel). The scavenging of reactive oxygen species (ROS) with N-acetylcyesteine (NAC) had no effect on IL-6 levels in EV cell line, ruling out the involvement of ROS production in this process (Fig. 5A and 5B). Thus, JNK activation contributed to the higher levels of IL-6 production in EVER2-deficient cells. This overproduction may involve a cooperative interaction between NF-κB and AP-1 sites in the IL-6 promoter or a JNK positive feedback amplification loop [33]. These findings demonstrated that PI3K and JNK activation was required to induce inflammation in EVER2-deficient cells, a process that was counteracted by an AKT-dependent signaling as shown by the increase of IL-6 production following treatment with Akt1/2 inhibitor (Fig. 5A).

We next investigated the kinases involved in the constitutive nuclear phosphorylation of c-jun in EVER2−/− cells. This phosphorylation was largely inhibited by Go6976, a PKCα inhibitor, and was completely abolished by SP60025, a JNK inhibitor (Fig. 5C). By contrast, no change was detected with ROS scavengers, such as NAC. Activation of the epidermal growth factor (EGF) receptor was not involved in the phosphorylation process, because specific inhibitors of this receptor did not prevent c-jun phosphorylation. Thus, EVER2 loss induced high basal levels of c-jun phosphorylation through a PKCα/JNK-dependent pathway.

The loss of EVER2 facilitates HPV5 LCR activation via the PI3K and JNK signaling pathways

The activation of AP-1 transcription factors, such as c-jun, is important for β-HPV transcription [34]. As basal levels of c-jun phosphorylation were higher in EVER2−/− cells than in control cells, we hypothesized that the AP-1 pathway might influence HPV5 LCR activation in EVER2−/− cells. We tested this hypothesis, by analyzing the expression of a luciferase reporter gene under the control of a full-length LCR from HPV5 (HPV5-ILCR/luc) containing AP-1-responsive elements. Luciferase activity levels were higher in the EV cell line than in controls (Fig. 5D). Similar results were obtained for the miEVER2 cell line (data not shown). The inhibition of PI3K and JNK by Ly294002 and SP60025, respectively, decreased luciferase activity more strongly in EVER2−/− cell lines than in controls (Fig. 5D). These data are consistent with constitutive activation of the JNK/AP-1 pathway in EVER2-deficient cells and show that EVER2 loss facilitates HPV5 LCR activation via the PI3K and JNK signaling pathways.

Overall, we demonstrated here that EVER2 loss promotes a PI3K- and JNK-dependent activation pathway, increasing HPV5 LCR activation and IL-6 production. By contrast, the NF-κB response to PMA+ionomycin was defective in EVER2−/− cells. We also found that EVER2-deficient cells responded poorly to TNF through impaired alternative and classical NF-κB pathways. All these findings suggest that EVER2 prevents PKCα-dependent JNK activation and targets TRAF2 through effects on its ubiquitination and localization, leading to efficient responses to TNF through the NF-κB and JNK/AP-1 signaling pathways.

Thus, our findings indicate that a genetic defect may create a microenvironment facilitating HPV transcription but preventing TNF through the NF-κB and JNK/AP-1 signaling pathways.

Materials and Methods

Cell lines and culture conditions

Keratinocytes were obtained from hair follicles from a healthy subject and from a Polish EV patient carrying a homozygous mutation characterized by a splice site mutation IS4-1G>T, T1506X3 mutation in the EVER2 gene [7] as described by Linnat A and F.K. Noser [35]. They were cultured on a feeder layer of lethally irradiated mouse 3T3-J2 fibroblasts [36]. Keratinocytes were immortalized with two lentiviruses encoding telomerase and SV40-AgT [37]. Immortalized cells were maintained in culture for four months in K-SFM medium (Gibco) without a feeder layer. The immortalized keratinocyte cell lines were then allowed to adapt to growth in serum-containing medium. The study was approved by the Ethics Committee at the Medical University of Warsaw. Written informed consent was received from all participants.

For EVER2 silencing, a microRNA expression vector specific for EVER2 (miEVER2) was generated with the BLOCK-iT™ Pol II miR RNAi Expression Vector Kit (Invitrogen) according to the manufacturer’s instructions. Two single-stranded DNA oligonucleotides encoding the pre-miRNA were designed with RNAi Designer (Invitrogen), according to the manufacturer’s instructions. These oligonucleotides targeted the following sequence of the EVER2 gene (encoding part of the N-terminal region of the protein): 3′-CAGGAATTCGTCCATTCTICA-3′. Cell lines from a healthy subject were transfected with miEVER2 or a control (miCTRL) expression vector containing an EozGFP-coding sequence in the presence of Lipofectamine 2000 reagent (Invitrogen) according to the manufacturer’s instructions. The cells expressing the pre-miEVER2 and control constructs were positive for green fluorescence protein (GFP) and were purified by cell sorting with a MoFLO flow cytometer (Beckman Coulter, Villepinte, France). All keratinocyte cell lines were grown in MEM containing 10% fetal calf serum, penicillin/streptomycin and 2 mM L-glutamin.

In some experiments, cells were transfected with pre-designed siRNAs (Applied Biosystems) directed against different parts (exons 5, 6, 8 or 10) of EVER2 (siEVER2) or control siRNAs (siCTRL). EVER2 transcripts levels were then assessed by semi-quantitative RT-PCR (Fig. 1B).

For all stimulation experiments, cells were serum-starved overnight and treated with 10 ng/ml recombinant human TNFα (TNF) or a combination of 10 ng/ml PMA plus 50 ng/ml ionomycin, for various periods of time. HEK-293T cells were cultured in DMEM containing 10% fetal calf serum, penicillin/streptomycin and 2 mM L-glutamin.

Antibodies and reagents

Antibodies against TRAF1, TRAF2, TRAF3, TRAF6, p50, p52, p105, p100, IkBz, the phosphorylated forms of IKKs (Ser176) β (Ser177), p65 (Ser536), c-jun (Ser63) and IL-6 were purchased from Cell Signaling Technology. Antibodies against p65 and TRAF5 were obtained from Santa Cruz Biotechnology. The antibody against EVER2 was purchased from Interchim. The anti-lamin B antibody was obtained from Thermo Scientific. Recombinant human TNFα was purchased from Miltenyi Biotec. Anti-actin, anti-Flag and anti-hemaglutinin (HA) antibodies, PMA, ionomycin, brefeldin A, Ly294002, SP60025, Bay 11-7082, Go6976, AG1478, Akt1/2 kinase inhibitor and N-acetyl-cysteine were obtained from Sigma-Aldrich. Anti-Ghuc and anti-GFp antibodies were obtained from BioLabs. Anti-KL1 antibody was purchased from Immunotech Biotechnologies. Anti-ubiquitin antibody was obtained from Dako.
Cytokine secretion

Cells were plated in 24-well plates and grown for 24 hours. They were serum-starved overnight and then maintained in culture for an additional 48 hours in the presence or absence of TNF. The supernatants were collected and the concentrations of IFN-γ, IL-1β, IL-2, IL-4, IL-5, IL-6, IL-8, IL-10 and IL-12 were determined by flow cytometry with the FlowCytomix human Th1/Th2 10plex bead kit and known standard curves, according to the manufacturer’s instructions (Bender MedSystems). Beads were analyzed with a Beckman Coulter EPICS-XL MCL flow cytometer (Coulter Electronics Inc., Hialeah, FL) and data were analyzed with Bender MedSystems software. All the experiments were performed in triplicate. Intracellular cytokine production was studied in cells plated in six-well plates and grown for 24 hours. The cells were serum-starved overnight and stimulated for 16 h with TNF in the presence of brefeldin A (5 μg/ml). They were then lysed for analysis by western blotting.

Transient transfection and reporter gene assays

The cells were plated in 24-well plates, grown to 50–80% confluence and transfected for 3 hours by the PEI (polycyteneamine) method, as previously reported [7]. For NF-κB luciferase reporter assays, we used 0.1 μg of NF-κB firefly luciferase reporter plasmid (Stratagene) per well. In some experiments, cells were transfected with the firefly luciferase reporter gene under the control of a LCR (nucleotides 189-37) from HPV5 (HPV5-LCR/luc) (0.3 μg per well) or a minimal region of the LCR (nucleotides 189-37) from HPV5 (HPV5-minLCR/luc) (0.3 μg per well). The Renilla luciferase plRlTK vector (0.03 μg) was used as internal control for transfection efficiency. Luciferase activity was assessed with the Dual-Luciferase Reporter Assay kit (Promega) and a Centro XS LB 960 luminometer (Berthold Technologies). Data were normalized against protein concentration, as determined by the Bio-Rad protein assay (Bio-Rad). All the experiments were performed in triplicate.

For transient EVER2 expression, cells were transfected with the EVER2 cDNA encoding the wild-type EVER2 protein. The full-length Flag-tagged EVER2 expression vector was obtained as previously reported [7]. Gluc2-tagged TRAF1, TRAF2 and RIPK1 expression vectors were constructed by inserting cDNAs encoding human TRAF1 or TRAF2 into pSPICA-N2 vector containing a Gaussia princeps luciferase fragment (Gluc2), as previously described [30].

For transient siRNA transfection experiments, cells were transfected in the presence of INTERFERin™ (Polyplus transfection), according to the manufacturer’s instructions.

Semi-quantitative and quantitative real-time PCR

Total RNA was isolated with Trizol ( Gibco BRL). cDNA was produced by reverse transcription with the Superscript III First-Strand system (Invitrogen). The oligonucleotide sequences used for the semi-quantitative real-time PCR (RT-PCR) were, for EVER2: 5’-CTCTTCCGCAAGAGATTCGCTGTG-3’ and 5’-GCTGTA-CACGGAGCTGCTGTCG-3’; and for GAPDH: 5’-GACCA-CAGTGGCATCCATACCT-3’ and 5’-TCCACACAACCCTGT-TGCTTAG-3’.

Cell fractionation and western blotting

Whole-cell extracts were prepared by incubating the cells for 45 minutes in lysis buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 0.1% SDS, 0.5% DOC, 1 mM EDTA, 1% NP40, 1× complete protease inhibitor cocktail, 0.1 mM Na3VO4 and 10 mM NaF, on ice. For some experiments, nuclear and cytosolic protein fractions were separated with NE-PER Nuclear and Cytoplasmic Extraction Reagents ( Pierce Biotechnology), according to the manufacturer’s instructions. For EVER2 detection, insoluble fractions were prepared as previously described [27]. In some experiments, Triton-soluble and -insoluble fractions were prepared as previously reported [28]. The various fractions were subjected to electrophoresis and the protein bands were transferred to nitrocellulose membranes, which were probed with the appropriate antibodies and developed with the SuperSignal West Pico or West Femto Chemiluminescent Substrate System (Pierce Biotechnology). Lamin B and β-actin were used as markers of the nuclear and cytoplasmic fractions, respectively. A Las-4000 image reader and Multi Gauge 3.1 software (Fuji Photo Film Co., Ltd.) were used for densitometric quantification of western blots. Data were normalized with respect to β-actin for the cytoplasmic fraction and lamin B for nuclear fractions.

Co-immunoprecipitation assay

Cells were lysed in immunoprecipitation buffer (50 mM Tris, pH 7.5, 150 mM NaCl, 1 mM EDTA, 0.5% NP40, 0.25% DOC, 1× complete protease inhibitor cocktail, 0.1 mM Na3VO4 and 10 mM NaF). Cell lysates were incubated overnight with 4 μg of antibody at 4°C. Complexes were precipitated with protein G/ protein A-agarose (Calbiochem), washed and suspended in SDS sample buffer. Immunoprecipitates were subjected to SDS-PAGE and western blotting.

Ubiquitination assay

For deubiquitination assays, HEK-293T cells were transfected with an HA-ubiquitin construct and other plasmids, as indicated. After 24 h, cells were stimulated with TNF for 2 h and harvested. Cell pellets were lysed as previously described [27]. The whole-cell extracts were subjected to immunoprecipitation and immunoblott- ing with the indicated antibodies. For endogenous TRAF2 ubiquitination assays, cells were plated and cultured for three days. They were then serum-starved for 6 h, harvested and lysed, as indicated above. Endogenous TRAF2 was immunoprecipitated with anti-TRAF2 antibody and subjected to immunoblotting with ubiquitin- and TRAF2-specific antibodies.

Immunostaining and microscopy

Adherent cells were fixed with methanol and incubated with anti-p65 (Santa Cruz), anti-TRAF2 (Santa Cruz) or anti-KL1 (Immunotech Biotechnologies) and then with the Alexa Fluor 488-conjugated F(ab’)2 fragment of anti-mouse IgG (Invitrogen). The slides were mounted in VECTASHIELD Mounting Medium with DAPI (Vector Laboratories). Images were acquired with a Nikon Eclipse 80i fluorescence microscope (Nikon) with a 20× objective, a 377/482 nm filter and a Dxm 1200C digital camera (Nikon). They were analyzed with NIS Elements Imaging software (Nikon).

The fluorescence intensity of nuclear p65 was quantified with ImageJ software (National Institutes of Health [NIH]).

Statistics

The statistical significance of differences was determined with Student’s t test. For all analyses, P<0.05 was considered significant.

Supporting Information

Figure S1 Schematic diagram illustrating the classical and alternative NF-κB signaling pathways, adapted from...
Classical pathway involves activation of IKK complex by IKK-mediated IkB phosphorylation, and subsequent degradation, resulting in nuclear translocation of the NF-κB heterodimer p65/p50. Alternative NF-κB pathway is dependent on NIK and IKKz and mediates the translocation of RelB/p52 complex. TNFR1 also activates JNK kinase, which activates AP-1 transcription factor.

Figure S2 NF-κB and c-jun activation in the EV cell line. EV and Healthy cell lines were left untreated or were stimulated with PMA+ionomycin (A) or TNF (B–E) for the times indicated. (A) Whole-cell lysates were analyzed by western blotting, to explore the IKKβ complex. The phosphorylated forms of IKKzβ are indicated as pIKKzβ. The IKKz and actin bands on western blots were quantified by densitometry. Results are reported as the ratio of IKKz to actin. The ratio of the Healthy cell line at time 0 was set to 1. (B–C) Cytoplasmic and nuclear fractions were analyzed for the classical NF-κB and AP-1 axis (B) or the alternative NF-κB axis (C). The phosphorylated forms of p65, and c-jun are indicated as p65 and p65-cJun, respectively. Whole-cell lysates (Total) were analyzed for total c-jun expression. The results shown are representative of three independent experiments. (D–E) Keratinocytes from the EV patient or the healthy subject were left untreated or were treated with TNF for 1 hour and stained for NF-κB (p65); the nucleus was stained with DAPI (D). Bars, 100 μm. (E) The fluorescence intensity of nuclear p65 was quantified before and after stimulation. Asterisks over error bars indicate statistically significant differences in fluorescence intensity between unstimulated and TNF-stimulated cells. Statistically significant differences between the two cell lines are indicated by asterisks over the brackets. Data are means ± SD of three independent experiments. *, P<0.05; **, P<0.01; ***, P<0.001.

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