The c-FLIP–NH₂ terminus (p22-FLIP) induces NF-κB activation

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c-FLIP proteins (isoforms: c-FLIP₁, c-FLIPₛ, and c-FLIPᵣ) play an essential role in the regulation of death receptor–induced apoptosis. Here, we demonstrate that the cytoplasmic NH₂-terminal pro-caspase-8 cleavage product of c-FLIP (p22-FLIP) found in nonapoptotic malignant cells, primary T and B cells, and mature dendritic cells (DCs) strongly induces nuclear factor κB (NF-κB) activity by interacting with the IκB kinase (IKK) complex via the IKKγ subunit. Thus, in addition to inhibiting apoptosis by binding to the death-inducing signaling complex, our data demonstrate a novel mechanism by which c-FLIP controls NF-κB activation and life/death decisions in lymphocytes and DCs.

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Abbreviations used: CHX, cycloheximide; DED, death effector domain; DISC, death-inducing signaling complex; EMSA, electrophoretic mobility shift assay; IKK, IκB kinase complex; siRNA, small interfering RNA.

c-FLIP is a well-described inhibitor of death receptor–mediated apoptosis (1). At the mRNA level, it can be found in multiple splice variants, whereas at the protein level only three isoforms, c-FLIP₁, c-FLIPₛ, and c-FLIPᵣ, have been detected so far (1–4). All three c-FLIP isoforms contain two death effector domains (DEDs), which are structurally similar to the NH₂-terminal part of procaspase-8. c-FLIP₁ also contains catalytically inactive caspase-like domains (p20 and p12).

c-FLIP proteins are recruited to the death-inducing signaling complex (DISC) by DED interactions (3–5). Both short c-FLIP isoforms, c-FLIPₛ and c-FLIPᵣ, block death receptor–induced apoptosis by inhibiting procaspase-8 activation at the DISC (2, 3). The role of c-FLIP₁ at the DISC is still a matter of controversy (6, 7). Some reports describe c-FLIP₁ as an antiapoptotic molecule, functioning in a way analogous to c-FLIPₛ, whereas others describe c-FLIP₁ as a proapoptotic molecule, facilitating the activation of procaspase-8 at the DISC. This proapoptotic role may explain the phenotype of c-FLIP₁–deficient mice characterized by heart failure and death at embryonic day 10.5. The same phenotype has been reported for caspase-8– and FADD–deficient mice (8–11).

In addition to its antiapoptotic role in death receptor–induced apoptosis, c-FLIP proteins were invoked to play a prominent role in NF-κB signaling (12–14). The transcription factor NF-κB family regulates the expression of genes crucial for innate and adaptive immune responses, cell growth, and apoptosis (15). In mammalian cells, the NF-κB family is composed of five members: RelA, RelB, c-Rel, p50/NF-κB1, and p52/NF-κB2 (16). In most cells, the NF-κB dimer is sequestered in the cytosol by inhibitors of the κB protein (IκB), and its nuclear translocation can be induced by a wide variety of stimuli (16). These stimuli trigger activation of the IκB kinase (IKK) complex, which consists of two catalytic subunits, IKKα and IKKβ, as well as a regulatory subunit, IKKγ/NEMO. When the IKK complex is activated, IκB is phosphorylated, and the IκBs are degraded in a ubiquitin-dependent manner. The NF-κB dimers can then be translocated into the nucleus, where target gene transcription is induced.

Recently, it has been demonstrated that overexpression of c-FLIP₁ activates NF-κB (13, 17). In another study, upon overexpression, c-FLIP₁ was shown to interact with established components of the TNFR-mediated NF-κB activation pathway, TRAF1, TRAF2, and RIP (12). In addition, it has been reported that c-FLIP₁–mediated NF-κB activation requires cleavage to p43-FLIP, also demonstrated to interact with TRAF2 (18). In TNFR-mediated NF-κB activation, TRAF2 and RIP were described to act upstream of the IKK complex (19, 20).

Here, we show that in nonapoptotic cells, c-FLIP forms heterodimers with procaspase-8 resulting in a novel NH₂-terminal fragment of c-FLIP (p22-FLIP). p22-FLIP turned out to be the key mediator of NF-κB activation by...
direct binding to the IKK complex. These findings provide a new mechanism of c-FLIP-mediated NF-κB activation and shed light on the regulation of life/death decisions made in lymphocytes.

RESULTS

A new form of c-FLIP can be detected in malignant B and T cells

In addition to the three previously described c-FLIP protein isoforms, c-FLIP<sub>L</sub>, c-FLIP<sub>R</sub>, and c-FLIP<sub>S</sub> (2, 3, 21), we have detected a new prominent protein band with the anti-FLIP mAb NF6 directed against the DED region of c-FLIP (Fig. 1 A). The molecular mass of this protein is ~22 kD. The p22 protein was observed in total cellular lysates (Fig. 1 A) and in immunoprecipitates (Fig. 1 B) from B lymphoblastoid cell lines Boe<sup>L</sup> and Raji and the T cell lines iHUT78 and Jurkat A3, but not in CEM and SKW6.4 cells. The viability of the cells used for analysis was verified by negative propidium iodide and annexin V staining (Fig. S1, available at http://www.jem.org/cgi/content/full/jem.20051556/DC1). P22 protein was the most prominent in Boe<sup>L</sup> cells (Fig. 1, A and B). We call this protein p22-FLIP.

The detection of p22-FLIP with the anti-FLIP mAb NF6 indicated the presence of DEDs in p22-FLIP because the antibody was raised against the NH<sub>2</sub>-terminus of c-FLIP. Furthermore, p22-FLIP disappeared upon the addition of zVAD-fmk (Fig. 1 C). This suggests that p22-FLIP is likely a caspase-dependent cleavage product of c-FLIP. We then analyzed the primary structure of c-FLIP<sub>L/S</sub> and found an aspartate residue at position 198 (Fig. 1 D). Cleavage at Asp<sup>198</sup> could result in the formation of an NH<sub>2</sub>-terminal DED-containing cleavage product with a molecular mass of ~22 kD, corresponding to p22-FLIP.

To test this hypothesis, we generated a cDNA corresponding to the c-FLIP NH<sub>2</sub>-terminal fragment resulting from cleavage at Asp<sup>198</sup>. Subsequently, p22-FLIP was translated in vitro and added to cell lysates of Boe<sup>L</sup> cells, followed by immunoprecipitation with the anti-FLIP mAb NF6. The products of immunoprecipitation as well as the corresponding lysates were analyzed by Western blot (Fig. 1 E). After adding the in vitro–translated c-FLIP NH<sub>2</sub>-terminal fragment, the band corresponding to p22-FLIP increased considerably. Thus, we conclude that the molecular mass of the in vitro–translated product was indeed identical to endogenous p22-FLIP. These data provide the first evidence that p22-FLIP is an NH<sub>2</sub>-terminal cleavage product of c-FLIP generated by cleavage at Asp<sup>198</sup>.

p22-FLIP identification as the NH<sub>2</sub>-terminal cleavage product of c-FLIP

To study whether p22-FLIP can be generated from both c-FLIP<sub>L</sub> and c-FLIP<sub>S</sub>, we performed the following experiment in vitro. c-FLIP<sub>L</sub>, c-FLIP<sub>S</sub>, and FLAG-c-FLIP<sub>L</sub> were translated in vitro, [35S] labeled, and added to total cellular lysates of HUT78 and J16 cells (Fig. 2 A). Upon incubation, all c-FLIPs were cleaved into the NH<sub>2</sub>-terminal fragment p22-FLIP (for c-FLIP<sub>L</sub> and c-FLIP<sub>S</sub>), the NH<sub>2</sub>-terminal FLAG-p22-FLIP (for FLAG-c-FLIP<sub>L</sub>), and the COOH-terminal fragment p33-FLIP (for c-FLIP<sub>L</sub> and FLAG-c-FLIP<sub>L</sub>). Consistent with the results obtained previously, p22-FLIP was not detected upon the addition of zVAD-fmk. Thus, we observed a caspase-dependent processing of c-FLIP<sub>L/S</sub> into p22-FLIP in nonapoptotic cells.

To provide conclusive evidence that p22-FLIP is indeed the cleavage product of c-FLIP resulting from cleavage at Asp<sup>198</sup>, we generated an uncleavable D198A mutant of FLAG-c-FLIP<sub>L</sub>. In vitro–translated [35S]-labeled D198A-FLAG-c-FLIP<sub>L</sub> and WT-FLAG-c-FLIP<sub>L</sub> were added to the lysates of HUT78 cells (Fig. 2 B). As anticipated, FLAG-p22-FLIP was generated in a caspase-dependent manner only from WT-FLAG-c-FLIP<sub>L</sub> into p22-FLIP in nonapoptotic cells.
construct (Fig. 2 C). Collectively, these data demonstrate that p22-FLIP is the NH₂-terminal cleavage product of c-FLIP generated by caspase cleavage at Asp198.

p22-FLIP is generated by procaspase-8 activity and inhibits death receptor–induced apoptosis

To unravel the mechanism of p22-FLIP formation and find the caspase directly involved in c-FLIP processing, we investigated whether procaspase-8 might generate p22-FLIP. Procaspase-8, which, as a proform, was reported to possess catalytic activity (7, 22) and form heterodimers with c-FLIP in the cytosol by DED interactions, represented a likely candidate (6, 23). Therefore, we immunoprecipitated procaspase-8 from HUT78 cells with an anti–caspase-8 mAb and added in vitro–translated [35S]-labeled c-FLIPL (Fig. 3 A, left). Interestingly, we observed cleavage of c-FLIPL into the NH₂-terminal fragment p22-FLIP and the COOH-terminal fragment p33-FLIP by procaspase-8. The processing was blocked by zVAD-fmk.

To ensure that the mature caspase-8 heterotetramer could not process c-FLIP into p22-FLIP, we added recombinant active caspase-8 to in vitro–translated c-FLIPL (Fig. 3 A, right). As expected, recombinant caspase-8 cleaves c-FLIPL in a “classic apoptotic” fashion and generated p43-FLIP and p12-FLIP. Thus, we show that caspase-8 and procaspase-8 process c-FLIP in two mutually exclusive ways. The active caspase-8 heterotetramer generates the well-characterized p43-FLIP and p12-FLIP cleavage products, whereas procaspase-8 activity induces formation of the novel p22-FLIP cleavage product as well as the COOH-terminal p33-FLIP. This is consistent with a report on different substrate specificities of the caspase-8 proform and active caspase-8 (7). Clearly, the present study is the first to demonstrate two different caspase-8 specificities with respect to c-FLIP cleavage.

c-FLIP proteins are well-established inhibitors of death receptor–mediated apoptosis. To examine the role of p22-FLIP in death receptor–mediated apoptosis, we generated stable BJAB cell lines overexpressing either high (p22-FLIPhigh) or low (p22-FLIPlow) amounts of p22-FLIP. The amount of p22-FLIP in these cell lines was validated by Western blot (Fig. 3, C and D, respectively). The reduction in apoptosis was more prominent in the p22-FLIPhigh cells. These data provide evidence that compared with other c-FLIP proteins, p22-FLIP is an even stronger inhibitor of death receptor–induced apoptosis.

c-FLIP proteins were reported to have a short half-life (24, 25). We and others have demonstrated that cycloheximide (CHX) sensitizes cells toward death receptor–induced apoptosis (Fig. 2).
Figure 3. p22-FLIP is generated by procaspase-8 and inhibits death receptor–induced apoptosis. (A) Procaspase-8 was immunoprecipitated from HUT78 cells using anti–caspase-8 mAb C15 and then incubated for 1 h at 37°C together with in vitro–translated [35S]–labeled c-FLIPΔ in the presence or absence of zVAD-fmk. c-FLIP processing was analyzed by autoradiography (top left). c-FLIP cleavage products p22 and p33 are indicated. Afterward, the same membrane was subjected to Western blot analysis using anti–caspase-8 mAb C15 (bottom left). [35S]–labeled c-FLIPΔ was incubated with the indicated concentrations of recombinant caspase-8 for 1 h at 37°C. c-FLIP processing was analyzed by autoradiography (top right). c-FLIP cleavage products p12 and p43 are indicated. Afterward, the same membrane was subjected to Western blot analysis using anti–caspase-8 mAb C15 (bottom right). (B) Analysis of p22-FLIP expression in BJAB cell lines stably overexpressing high or low amounts of p22-FLIP (p22-FLIPhigh or p22-FLIPlow, respectively). Endogenous expression of c-FLIP Δ is used as a loading control. (C) p22-FLIPhigh, p22-FLIPlow, and vector-transfected BJABs (Ctrl.) were stimulated with 1 μg/ml anti–APO-1 antibodies or 50 μl/ml LZ-CD95L for 16 h. Specific cell death was calculated as described in Materials and methods. (D) p22-FLIPhigh, p22-FLIPlow, and
apoptosis, which correlates with a decrease of c-FLIP levels (24, 25). To understand whether sensitization also involves a decrease of the p22-FLIP level, we studied Boe cells that contain high levels of p22-FLIP. Treatment of Boe cells with CHX resulted in a substantial decrease of the p22-FLIP level within 4 h (Fig. S2 A, available at http://www.jem.org/cgi/content/full/jem.20051556/DC1). This decrease correlated with an enhanced sensitivity toward CD95- and TRAIL-induced apoptosis (Fig. S2 B). These data provide additional evidence that the p22-FLIP level in Boe cells correlates with the sensitivity toward death receptor–induced apoptosis, pointing toward the inhibitory role of p22-FLIP.

The antiapoptotic action of c-FLIP in death receptor–mediated apoptosis involves inhibition of caspase-8 activation at the DISC (2, 3, 24). To investigate whether p22-FLIP is also recruited to the DISC, we immunoprecipitated the CD95 DISC from p22-FLIP<sub>high</sub> cells. Indeed, p22-FLIP was recruited to the DISC (Fig. 3 E). As expected, the activation of procaspase-8 at the DISC of p22-FLIP<sub>high</sub> cells was lower than in vector-transfected cells. The amount of caspase-8 cleavage products in the DISC, p43/p41, and p18 was markedly reduced in p22-FLIP<sub>high</sub> cells, whereas the amounts of FADD and CD95 were similar in both p22-FLIP<sub>high</sub> and vector-transfected cells. This observation is consistent with the reduced sensitivity of p22-FLIP<sub>high</sub> cells toward CD95-induced apoptosis and shows that p22-FLIP effectively blocks caspase-8 activation at the DISC, thereby inhibiting CD95-induced apoptosis.

Figure 4. p22-FLIP is a strong inducer of NF-κB. (A) 293T cells were cotransfected with MEKK1, p22-FLIP, c-FLIP<sub>l</sub>, and luciferase reporter plasmid. GFP transfections were performed to control the transfection efficiency. Western blot analysis using anti-FLIP mAb NF6 was performed to control equal protein expression (right). (B) 293T cells were cotransfected with p22-FLIP and the luciferase reporter plasmid. After the indicated periods of time, cells were lysed and NF-κB luciferase activity was determined (top). Western blot analysis using anti-FLIP mAb NF6 was performed to determine the expression level of p22-FLIP. (C) Nuclear extracts, which were prepared from 293T cells transfected with p22-FLIP or GFP, were subjected to EMSAs using 32P-labeled oligonucleotides containing an NF-κB (left) or an NF-Y (right) binding site. p22-FLIP expression was verified by Western blot.

vector-transfected BJABs (Ctrl.) were stimulated with the indicated concentrations of FLAG-TRAIL for 16 h. (E) CD95 DISCs were immunoprecipitated from 5 × 10<sup>7</sup> cells of p22-FLIP<sub>high</sub> and vector-transfected BJABs (Ctrl.) and analyzed by Western blot with anti–caspase-8 mAb C15, anti-FLIP mAb NF6, anti-CD95 polyclonal antibody C20, and anti-FADD mAb.
p22-FLIP is a strong inducer of NF-κB

We established an inhibitory role of p22-FLIP in death receptor–induced apoptosis; however, the questions of why p22-FLIP is present in nonapoptotic malignant cells and which functional role p22-FLIP might play in the non-apoptotic scenario were not answered. Using NF-κB luciferase activation assays with p22-FLIP and c-FLIPL, we observed that p22-FLIP is a strong inducer of NF-κB (Fig. 4A). Moreover, p22-FLIP–mediated NF-κB activation was much stronger than that observed with c-FLIPL, even though expression levels of p22-FLIP and c-FLIPL were similar (Fig. 4A).

To clarify whether NF-κB activation correlates with the expression level of p22-FLIP, we performed transient transfections of 293T cells with p22-FLIP (Fig. 4B). The maximum of NF-κB activity at 10 h paralleled the increase of p22-FLIP expression. Moreover, the observed expression level of p22-FLIP in 293T cells (Fig. 4B) compared with c-FLIPL was still lower as compared with the ratio p22-FLIP/c-FLIPL in BoeR cells (Fig. 1A). Thus, we demonstrate that the induction of NF-κB is specific for p22-FLIP and does not depend on a high expression level of p22-FLIP.

We also assayed p22-induced NF-κB activity by electrophoretic mobility shift assay (EMSA; Fig. 4C), which independently confirmed that p22-FLIP induces NF-κB. Thus, we demonstrated that p22-FLIP is a strong inducer of NF-κB.

p22-FLIP induces NF-κB by direct interaction with the IKK complex

To get more insight into the mechanism of p22-mediated NF-κB induction, we coexpressed p22-FLIP with the inhibitors of NF-κB (IkBa and IkBβ) and with components of the IKK complex (IKKα, IKKβ, DN-IKKα, and DN-IKKβ; Fig. 5A). Cotransfections of p22-FLIP with increasing amounts of IkBa and IkBβ inhibited NF-κB activation. Looking at more upstream events, we observed that cotransfection with DN-IKKα and DN-IKKβ also led to suppression of NF-κB activation. Collectively, the results indicated that p22-FLIP is a strong inducer of NF-κB acting via the canonical NF-κB pathway.

We further examined whether p22-FLIP directly interacts with the IKK complex. FLAG-tagged IKKα, IKKβ, and IKKγ were transiently cotransfected with p22-FLIP into 293T cells, and then immunoprecipitated with anti-FLAG and anti-FLIP antibodies (Fig. 5B). We did not observe any interaction between p22-FLIP and IKKα or IKKβ. However, p22-FLIP was coimmunoprecipitated with FLAG-IKKγ and vice versa. Thus, we showed that p22-FLIP interacts with the IKK complex via IKKγ.

In addition, we compared NF-κB induction by p22-FLIP, c-FLIPL, and p43-FLIP upon transient transfections in 293T cells (Fig. 5C). Interestingly, the addition of zVAD-fmk resulted in a decrease in NF-κB activation for p43-FLIP and c-FLIPL, but did not affect p22-FLIP–mediated NF-κB induction. These results strongly indicate that p43-FLIP and c-FLIPL require further cleavage to induce NF-κB activity, whereas p22-FLIP does not. Thus, we show that for the induction of NF-κB activity, both p43-FLIP and c-FLIPL have to be processed into p22-FLIP.

Figure 5. p22-FLIP induces NF-κB by direct interaction with the IKK complex. (A) 293T cells were cotransfected with luciferase reporter plasmid and either MEKK1, p22-FLIP, or c-FLIP, (top part of the diagram). 293T cells were cotransfected with p22-FLIP, the luciferase reporter plasmid, and any one of the constructs IkBa, IkBβ, WT-IKKα, WT-IKKβ, mutated IKKα, or IKKβ (bottom part of the diagram). Transfection efficiency was examined using GFP transfections. NF-κB luciferase activity was determined as described in Materials and methods. (B) FLAG or FLIP immunoprecipitations were performed from 293T cells that were transfected with p22-FLIP and any one of the constructs FLAG-IKKα, FLAG-IKKβ, or FLAG-IKKγ. Immunoprecipitated products were subjected to 12% SDS-PAGE gels and analyzed by Western blot using anti-FLAG mAb NF6 and anti-FLAG mAb. (C) 293T cells were cotransfected with MEKK1, p22-FLIP, c-FLIP, p43-FLIP, and the luciferase reporter plasmid. Transfected cells were incubated for 16 h in the presence of the indicated concentrations of zVAD-fmk and lysed, and NF-κB luciferase activity was determined.
p22-FLIP induces NF-κB during activation of primary lymphocytes and maturation of primary DCs

Next, we examined primary human T and B cells for the presence of p22-FLIP. Interestingly, p22-FLIP was absent in freshly prepared cells, but was generated upon activation of these cells with PHA (Fig. 6 A). The same phenomenon was observed in DCs. p22-FLIP was generated upon LPS stimulation, indicating that p22-FLIP is present during maturation of DCs (Fig. 6 A). Also, the increase of p22-FLIP levels in primary cells correlated with the increase of c-FLIP levels and, correspondingly, with the increase of the ratio of c-FLIP to pro-caspase-8. This observation provides additional evidence for the proposed mechanism of pro-caspase-8–mediated c-FLIP cleavage to p22-FLIP (Fig. 3).

To find out whether p22-FLIP also directly interacts with the IKK complex in primary human T cells, we performed immunoprecipitations using an anti-IKKγ antibody (Fig. 6 B). We observed p22-FLIP in the IKK complex. Thus, we conclude that p22-mediated NF-κB induction occurs via the same mechanism in primary cells.

To obtain more insight into the role of p22-FLIP in T cell activation, we studied the proliferation of primary T cells upon silencing of c-FLIP using small interfering RNA (siRNA). The silencing of c-FLIP resulted in down-regulation of c-FLIPL/S/R as well as its cleavage product, p22-FLIP (Fig. 6 C). Primary T cells were stimulated with either PHA or anti-CD3/CD28, and thymidine incorporation was measured after 3 d of additional culture (Fig. 6 C). Interestingly,
silencing of c-FLIP led to a complete stop of cell proliferation. Thus, we show that the absence of c-FLIP and, consequently, p22-FLIP led to severe defects in T cell proliferation.

**DISCUSSION**

c-FLIP proteins were demonstrated to induce NF-κB activation (12–14). However, the exact underlying mechanism of this process has not been established yet. In this study, we identified a new mechanism of c-FLIP–mediated NF-κB activation and showed that NF-κB activation requires c-FLIP processing into the NH2-terminal DED-containing fragment, p22-FLIP (Fig. 7). p22-FLIP is generated by procaspase-8 cleavage of both c-FLIP isoforms, c-FLIP.L and c-FLIP.S. Furthermore, p22-FLIP is a strong activator of NF-κB, acting directly at the level of the IKK complex by binding to IKKγ. In addition to its role as an activator of NF-κB, p22-FLIP can block apoptosis by directly binding to the DISC.

p22-FLIP is generated by cleavage of c-FLIP.L at Asp198. The resulting NH2-terminal fragment, p22-FLIP, contains two tandem DEDs and has a high structural homology to v-FLIP of the Kaposi’s sarcoma-associated herpesvirus and other v-FLIPs (14) that are also characterized by the presence of two DEDs followed by a short COOH terminus.

We argue that p22-FLIP is the final cleavage product of c-FLIP that serves as a mediator of NF-κB activation. This result is in contrast to previous studies, where it was suggested that p43-FLIP is a cleavage product of c-FLIP that mediates NF-κB activation (18). We have clearly shown that upon zVAD-fmk treatment, p43-FLIP–mediated activation of NF-κB is decreased, whereas the p22-FLIP–mediated NF-κB activation (12–14). However, the exact underlying mechanism of this process has not been established yet. In this study, we identified a new mechanism of c-FLIP–mediated NF-κB activation and showed that NF-κB activation requires c-FLIP processing into the NH2-terminal DED-containing fragment, p22-FLIP (Fig. 7). p22-FLIP is generated by procaspase-8 cleavage of both c-FLIP isoforms, c-FLIP.L and c-FLIP.S. Furthermore, p22-FLIP is a strong activator of NF-κB, acting directly at the level of the IKK complex by binding to IKKγ. In addition to its role as an activator of NF-κB, p22-FLIP can block apoptosis by directly binding to the DISC.

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In addition, p22-FLIP can block death receptor–mediated apoptosis by binding to the DISC via DED interactions and inhibiting procaspase-8 activity (left side).

Figure 7. The dual function of p22-FLIP in the cell. P22-FLIP induces NF-κB by interacting with IKKγ in the IKK complex (right side). In addition, p22-FLIP can block death receptor–mediated apoptosis by binding to the DISC via DED interactions and inhibiting procaspase-8 activity (left side).

Importantly, our study demonstrates a new NF-κB–activating pathway initiated by procaspase-8. We show that independently of death receptor stimulation, procaspase-8 generates the p22-FLIP cleavage product, which leads to the induction of NF-κB (Fig. 7). Recently, procaspase-8 was reported to play a prominent role in NF-κB induction via its involvement in the MALT1–Bcl-10 adaptor complex that is formed upon TCR stimulation (26). Our findings further elucidate the involvement of procaspase-8 in NF-κB induction.

Interestingly, we could also show that the active mature caspase-8 heterotetramer p102-p182 cleaves c-FLIP in vitro to p43-FLIP and p12-FLIP, but not to p22-FLIP. Thus, we observe that procaspase-8 and caspase-8 heterotetramer cleave c-FLIP in two different ways. This is consistent with reports upon different substrate specificities of the caspase–8 proform and active caspase-8 (7). Of note, our study is the first one demonstrating two different caspase-8 specificities with respect to c-FLIP cleavage.

In addition, we have observed in primary cells that upon increase of the ratio of c-FLIP to procaspase-8, the amount of p22-FLIP was substantially increased. It is likely that procaspase-8 constitutively cleaves c-FLIP to p22-FLIP, forming dimers with c-FLIP. The formation of such dimers between procaspase-8 and c-FLIP in the cytosol was described previously (6, 23). Thus, the ratio of procaspase-8 to c-FLIP in a particular cell type would be the crucial factor defining the amount of generated p22-FLIP and, correspondingly, the potential to induce NF-κB.

Our findings provide a molecular mechanism for how c-FLIP and procaspase–8 contribute to the activation and proliferation of primary lymphocytes. It was reported that caspase activity is essential for T cell activation, as it was shown in experiments with caspase inhibitors (27, 28). Analysis of caspase-8 and c-FLIP conditional knockout mice has demonstrated that those mice show severe defects in T cell activation and proliferation (29). Caspase-8 was reported to be essential for antigen-induced NF-κB activation in T, B, and NK cells (26, 30, 31). We show that upon silencing of c-FLIP in primary cells, the proliferation of these cells is impaired. This demonstrates the importance of our new p22-FLIP–mediated NF-κB–activating pathway in primary lymphocytes.

In conclusion, we described a new NF-κB–activating pathway, which is mediated by two well-described apoptotic DED-containing proteins: procaspase-8 and c-FLIP via their cleavage product, p22-FLIP. The balance between DED-containing proteins may provide sensitive signaling check points that cells use for signaling cross-talk and switching between apoptosis-resistant and -sensitive phenotypes and, thus, between life and death.

**MATERIALS AND METHODS**

**Cell lines.** The T cell lines HUT78, CEM, H9, Jurkat (clone A3), and Jurkat (clone J16); the B lymphoblastoid cell lines SKW6.4, Raji, and BJAB; and the pre-B cell line Boe were maintained in RPMI 1640, 10 mM Hepes, 50 μg/ml gentamycin, and 10% fetal calf serum (all from Life Technologies) in 5% CO2.
Antibodies and reagents. Anti-FADD mAb (IgG1) was purchased from Transduction Laboratories. Anti-FLAG mAb was purchased from Sigma-Aldrich. Anti-CD95 polyclonal antibody C20 was purchased from Santa Cruz Biotechnology, Inc. Anti-caspase-8 mAb C15 (mouse IgG2b) recognizes the p18 subunit of caspase-8 (32). Anti-IKKα/β and anti-IKKγ antibodies were purchased from Santa Cruz Biotechnology, Inc., and anti-ERK1 mAb was purchased from BD Biosciences. Anti–APO-1 is an agonistic mAb (IgG3, κ) recognizing an epitope on the extracellular part of CD95 (APO-1/Fas; reference 33). FLAG-TRAIL was obtained from H. Walczak (DKFZ, Heidelberg, Germany). Horseradish peroxidase– conjugated goat anti–mouse IgG1, IgG2a, and IgG2b were from SouthernBioTech. All chemicals used were of analytical grade and purchased from Merck or Sigma-Aldrich. Plas- mids encoding c-FLIPL and c-FLIP were prepared by bp cloning. The plasmid encoding FLAG–c-FLIPL has also been described (3). NF-kB reporter plasmid was provided by M. Li-Weber (DKFZ, Heidelberg, Germany). Constructs of FLAG-IKKe, FLAG-IKKβ, DN-FLAG-IKKe (K44A), and DN-FLAG-IKKβ (K44A) were provided by H. Nakano (Juntendo University, Tokyo, Japan).

EMSA. Soluble nuclear proteins were prepared and used for EMSA as described previously (34). For each reaction, 10–20 fmol of 32P-labeled oligonucleotides comprising an NF-κB binding site (5′-TCAGAGGGGAC-TTTCGAGAGGC-3′) or NF-Y binding site (5′-CACCCTTTTACCA-TATCAGAAAAAT-3′) were used.

Cloning of p22-FLIP and D198A-FLAG-c-FLIP L. p22-FLIP was cloned into the pEF4 expression vector (Invitrogen) using the PCR and the primer sequences: sense (encoding the KpnI restriction site): 5′-ggggatcc-ATGTCCTGTGAAGTCACTCC-3′ and antisense (encoding the XbaI restriction site): 5′-gttcagagctACTCTGAGACTCTTGG-3′.

The D198A-FLAG-c-FLIP L mutant was cloned via overlap-PCR into the pEF4 expression vector (Invitrogen). The KpnI–XbaI part was amplified using the following: forward primer 1: 5′-ggggatccc-ATCTTGTGTGAAGTCACTCC; reverse primer 1: 5′-CCCTGAAATTTGAGTGCCTGACTCTTGG-3′; reverse primer 2: 5′-CCCTGAAATTTGAGTGCCTGACTCTTGG-3′; forward primer 2: 5′-CCATAGGAAGCTGCTACTACAACTTTACAA-3′. The overlap-PCR was performed using the forward and reverse primer 2.

CD95 DISC analysis. Composition of the CD95 DISC was determined as follows. 5 × 105 cells were treated either with 1 μg/ml anti–APO-1 (IgG3) for 5 min at 37°C or left untreated, washed twice in 1× PBS, and lysed in lysis buffer (30 mM Tris/HCl, pH 7.5, 150 mM NaCl, 2 mM EDTA, 1 mM phenylmethylsulfonyl fluoride [Sigma-Aldrich], protease inhibitor cocktail [Roche], 1% Triton X-100 [Serva], and 10% glycerol). If pretreated with zVAD-fmk, cells were preincubated for 30 min at 37°C with the indicated concentrations of zVAD-fmk before stimulation. CD95 was immunoprecipitated with anti–APO-1 and protein A–Sepharose beads (Sigma-Aldrich) overnight at 4°C. Beads were washed five times with 20 volumes of lysis buffer. Immunoprecipitates were used for in vitro or in situ assay or analyzed using SDS-PAGE gels (35). Gels were transferred to Hybond nitrocellulose membrane (GE Healthcare), blocked with 5% nonfat dry milk in PBS/Tween (PBS plus 0.05% Tween 20) for 1 h, washed with PBS/Tween, and incubated with the primary antibody in PBS/Tween overnight at 4°C. Blots were developed with a chemoluminescence method according to the manufacturer’s protocol (PerkinElmer).

Immunoprecipitations. For c-FLIP immunoprecipitation, 5 × 105 cells were lysed in a volume of 1 ml for 30 min at 0°C, followed by the addition of 100 μl NF6 hybridoma supernatant together with 30 μl protein A–Sepharose. For FLAG immunoprecipitation, 2 × 106 cells were transfected using the calcium phosphate method 1 d before lysis. Immunoprecipitation was performed by using 4 μg anti-FLAG mAb together with 30 μl protein A–Sepharose. For IKKγ immunoprecipitation, 103 primary human T cells were lysed with or without stimulation, and the immunoprecipitation was performed by using 2 μg anti-IKKγ mAb together with 30 μl protein A–Sepharose. Immunoprecipitations were performed for 1 h at room temperature or overnight at 4°C. Beads were then washed five times with 20 volumes of lysis buffer and subjected to Western blot analysis as described above.

In vitro c-FLIP cleavage assays. Lysates of the indicated cell lines were prepared as described above and incubated with in vitro–translated [35S]-labeled c-FLIP L, FLAG-c-FLIP L, D198A-FLAG-c-FLIP L, or c-FLIP L (TNT, T7–coupled reticulocyte lysate system; Promega) overnight at 4°C (36). The procaspase-8 cleavage assay was performed as follows: 5 × 107 HUT78 cells were lysed, and procaspase-8 was immunoprecipitated using 100 μl anti–cas- passase-8 C15 hybridoma together with 30 μl protein A–Sepharose. Beads with bound procaspase-8 were incubated in reaction buffer (50 mM Heps, pH 7.4, 100 mM NaCl, 0.1% CHAPS, 10 mM dithiothreitol, and 10% sucrose) for 1 h at 37°C together with in vitro–translated c-FLIP L. The recombinaat procaspase-8 cleavage assay was performed as follows: 5 or 50 ng/ml recombinant caspase-8 was incubated in reaction buffer for 1 h at 37°C together with in vitro–translated c-FLIP L. Reactions were separated on 12% SDS-PAGE gels, blotted, and subjected to autoradiography.

Transfection of BJAB and Boe® cells. Stable transfection of BJAB as well as transient transfection of Boe® cells was performed using expression vectors or the empty vector by electroporation (950 μF, 200 V). Selection pressure was added 48 h after transfection (100 μg/ml zeocin) for 2 wk (BJAB cells) or 10 d (Boe® cells). Expression was controlled by Western blot using anti-FLIP mAb NF6. The p22-FLIP–expressing BJABs as well as the empty vector–transfected control cells were subcloned.

Cytotoxicity assay. To assay apoptosis, 5 × 105 cells were incubated in 48-well plates with or without the indicated amounts of anti–APO-1, LZ-CD95L, or FLAG-TRAIL for 16 h at 37°C. Cell death was measured by FSC/SSC via flow cytometry, and specific cell death was calculated as follows: (percentage of experimental cell death − percentage of spontaneous cell death)/100 × percentage of spontaneous cell death) × 100.

NF-κB activation assay. The day before transfection, 24-well titer plates were seeded with 0.5 × 105 293T cells. The cells were transfected using the calcium phosphate method with various expression vectors together with 500 ng of the NF-κB–driven luciferase reporter plasmid. Cells were washed with PBS 16 h after transfection and lysed for 20 min at room temperature in 50 μl lysis buffer (passive lysis buffer; Promega), followed by centrifugation (10,000 g) for 20 min to sediment insoluble materials. A total of 5 μl of cell lysates was mixed with 50 μl of the luciferase assay mixture (470 μM Beetle Luciferin [Promega], 1.07 mM (MgCO3)4Mg(OH)2 × 5 H2O, 20 mM N-Tris(hydroxymethyl)methylglycine, 2.67 mM MgSO4, 100 μM EDTA, 33.3 mM DTT, 270 μM CoA(OAc), and 530 μM ATP), and relative light units were measured with a Berthold dualoluminomat (Bad Wildbad).

Preparation and activation of primary human lymphocytes and DCs. Human peripheral T and B cells were prepared as described previously (37). For activation, resting primary human T cells (day 0) were cultured at 2 × 106 cells/ml with 1 μg/ml PHA (for 16 h (day 1), and primary human B cells (day 0) were cultured at 2 × 106 cells/ml with 1 μg/ml PHA. After preparing lymphocytes, the primary human monocytes were isolated using cell adhesion onto cell culture flasks. Leukocytes were resuspended in 20–30 ml RPMI 1640 with 10% FCS, and 2-ml aliquots were seeded into six-well titer plates. After incubation for 1 h, adherent cells were washed with PBS. Monocytes were differentiated into immature DCs by adding 1% human AB serum, 1% donor plasma, 1,000 U/ml GM-CSF (Schering), and 500 U/ml IL-4 (Immunotools) for 3 d. Cytokines were renewed after 3 d for an additional 3 d, and immature DCs were stimulated with 500 ng/ml LPS for 16 h.
CHX experiments. For CHX treatments, cells were incubated with 10 μg/ml CHX for the indicated periods of time. For assaying apoptosis in a cytotoxicity assay, 10^6 cells were pretreated with 10 μg/ml CHX for 4 h, washed, and incubated with the indicated concentrations of 1 μg/ml anti–APO-1 or 50 ng/ml FLAG-TRAIL for 16 h at 37°C in 24-well plates. Cell death was measured by FSC/SSC via flow cytometry, and specific cell death was calculated as follows: (percentage of experimental cell death – percentage of spontaneous cell death)/100.

Annexin V and propidium iodide staining. To detect phosphatidylserine exposure by flow cytometry, the T and B cell lines were washed once with PBS, incubated for 10 min on ice in 400 μl annexin V–FITC (Qbiogene) or 10 μg/ml propidium iodide (Invitrogen), and analyzed via flow cytometry.

siRNA-mediated knockdown of c-FLIP and proliferation assays. Primary human T cells were transfected by HiPerfect (QIAGEN) with siRNA-mediated knockdown of c-FLIP and proliferation assays. 10^5 cells were seeded into a 96-well titer plate and stimulated with 1 μg/ml anti–αCD3/αCD28 for 4 d. Proliferation was measured with a scintillation counter after tritiated thymidine ([3H]TdR) incorporation during the final 15–18 h of the culture.

Online supplemental material. Fig. S1 shows the living status of tumor cell lines used, and Fig. S2 shows that the resistance of BoeR cells toward CD95- or TRAIL-induced apoptosis is mediated by c-FLIP. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20051556/DC1.

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REFERENCES

1. Krüger, A., S. Baumann, P.H. Krammer, and S. Kirchhoff. 2001. FLICE-inhibitory proteins: regulators of death receptor-mediated apoptosis. Mol. Cell. Biol. 21:8247–8254.
2. Golks, A., D. Brenner, C. Fritsch, P.H. Krammer, and I.N. Lavrik. 2002. The long form of FLIP ablates cell death induction by the TNF receptors, Fas/Apo1, and DR3 and is lethal prenatally. Immunity. 9:267–276.
3. Kataoka, T., R.C. Budd, N. Hollier, M. Thome, F. Martinson, M. Irmler, K. Burns, M. Hahne, N. Kennedy, M. Kovacsovic, and J. Tschopp. 2000. The caspase-8 inhibitor FLIP promotes activation of NF-κB and Erk signaling pathways. Curr. Biol. 10:640–648.
4. Hur, W.H., H. Johnson, and H.B. Shu. 2000. Activation of NF-κB by FADD, Casper, and caspase-8. J. Biol. Chem. 275:10838–10844.
5. Thome, M., and J. Tschopp. 2001. Regulation of lymphocyte proliferation and death by FLIP. Nat. Rev. Immunol. 1:50–58.
6. Ghosh, S., and M. Karin. 2002. Missing pieces in the NF-κB puzzle. Cell. 109:S81–S96.
7. Hayden, M.S., and S. Ghosh. 2004. Signaling to NF-κB. Genes Dev. 18:2195–2224.
8. Chaudhary, P.M., M.T. Eby, A. Jasmin, A. Kumar, L. Liu, and L. Hood. 2000. Activation of the NF-κB pathway by caspase 8 and its homologs. Oncogene. 19:4451–4460.
9. Kataoka, T., and J. Tschopp. 2004. N-terminal fragment of c-FLIP(L) processed by caspase 8 specifically interacts with TRAF2 and induces activation of the NF-κB signaling pathway. Mol. Cell. Biol. 24:2627–2636.
10. Hur, G.M., J. Lewis, Q. Yang, Y. Lin, H. Nakano, S. Nedospasov, and Z.G. Liu. 2003. The death domain kinase RIP has an essential role in DNA damage-induced NF-κB B activation. Genes Dev. 17:873–882.
11. Liu, Z.G. 2005. Molecular mechanism of TNF signaling and beyond. Cell Res. 15:24–27.
12. Peter, M.E., and P.H. Krammer. 2003. The CD95(APO-1/Fas) DISC and beyond. Cell Death Differ. 10:26–35.
13. Boatright, K.M., M. Renatus, F.L. Scott, S. Sperandio, H. Shin, I.M. Pedersen, J.E. Ricci, W.A. Edrs, D.P. Sutherland, D.R. Green, and G.S. Salvesen. 2003. A unified model for apical caspase activation. Mol. Cell. 11:529–541.
14. Boatright, K.M., C. Denis, J.B. Denault, D.P. Sutherland, and G.S. Salvesen. 2004. Activation of caspases-8 and -10 by FLIP(L). Biochem. J. 382:651–657.
15. Schmitz, I., H. Weyd, A. Krüger, S. Baumann, S.C. Fas, P.H. Krammer, and S. Kirchhoff. 2004. Resistance of short term activated T cells to CD95-mediated apoptosis correlates with de novo protein synthesis of c-FLIPshort. J. Immunol. 172:2194–2200.
16. Fulda, S., E. Meyer, and K.M. Debatin. 2000. Metabolic inhibitors sensitize for CD95 (APO-1/Fas)-induced apoptosis by down-regulating Fas-associated death domain-like interleukin 1-converting enzyme inhibitory protein expression. Cancer Res. 60:3947–3956.
17. Su, H., N. Bidere, L. Zheng, A. Cubre, K. Sakai, J. Dale, L. Salmena, R. Hakem, S. Strauss, and M. Lenardo. 2005. Requirement is an activator of caspase-8 at the Fas death-inducing signaling complex. Biol. Chem. 277:45162–45171.
18. Chang, D.W., Z. Xing, V.L. Capacio, M.E. Peter, and X. Yang. 2003. Interdigimer processing mechanism of procaspase-8 activation. EMBO J. 22:4132–4142.
19. Kataoka, T., R.C. Budd, N. Hollier, M. Thome, F. Martinson, M. Irmler, K. Burns, M. Hahne, N. Kennedy, M. Kovacsovic, and J. Tschopp. 2000. The caspase-8 inhibitor FLIP promotes activation of NF-κB and Erk signaling pathways. Curr. Biol. 10:640–648.
20. Kataoka, T., and J. Tschopp. 2004. N-terminal fragment of c-FLIP(L) processed by caspase 8 specifically interacts with TRAF2 and induces activation of the NF-κB signaling pathway. Mol. Cell. Biol. 24:2627–2636.
21. Kataoka, T., and J. Tschopp. 2004. N-terminal fragment of c-FLIP(L) processed by caspase 8 specifically interacts with TRAF2 and induces activation of the NF-κB signaling pathway. Mol. Cell. Biol. 24:2627–2636.
22. Kataoka, T., and J. Tschopp. 2004. N-terminal fragment of c-FLIP(L) processed by caspase 8 specifically interacts with TRAF2 and induces activation of the NF-κB signaling pathway. Mol. Cell. Biol. 24:2627–2636.
23. Kataoka, T., and J. Tschopp. 2004. N-terminal fragment of c-FLIP(L) processed by caspase 8 specifically interacts with TRAF2 and induces activation of the NF-κB signaling pathway. Mol. Cell. Biol. 24:2627–2636.
for caspase-8 in NF-kappaB activation by antigen receptor. Science. 307:1465–1468.

27. Kennedy, N.J., T. Kataoka, J. Tschopp, and R.C. Budd. 1999. Caspase activation is required for T cell proliferation. J. Exp. Med. 190:1891–1896.

28. Misra, R.S., D.M. Jelley-Gibbs, J.Q. Russell, G. Huston, S.L. Swain, and R.C. Budd. 2005. Effector CD4+ T cells generate intermediate caspase activity and cleavage of caspase-8 substrates. J. Immunol. 174:3999–4009.

29. Chau, H., V. Wong, N.J. Chen, H.L. Huang, W.J. Lin, C. Mirtsos, A.R. Elford, M. Bonnard, A. Wakeham, A.I. You-Ten, et al. 2005. Cellular FLICE-inhibitory protein is required for T cell survival and cycling. J. Exp. Med. 202:405–413.

30. Chun, H.J., L. Zheng, M. Ahmad, J. Wang, C.K. Speirs, R.M. Siegel, J.K. Dale, J. Puck, J. Davis, C.G. Hall, et al. 2002. Pleiotropic defects in lymphocyte activation caused by caspase-8 mutations lead to human immunodeficiency. Nature. 419:395–399.

31. Dohrman, A., T. Kataoka, S. Cuenin, J.Q. Russell, J. Tschopp, and R.C. Budd. 2005. Cellular FLIP (long form) regulates CD8+ T cell activation through caspase-8-dependent NF-kappaB activation. J. Immunol. 174:5270–5278.

32. Scaffidi, C., J.P. Medema, P.H. Krammer, and M.E. Peter. 1997. FLICE is predominantly expressed as two functionally active isoforms, caspase-8/a and caspase-8/b. J. Biol. Chem. 272:26953–26958.

33. Trauth, B.C., C. Klas, A.M. Peters, S. Matzku, P. Moller, W. Falk, K.M. Debatin, and P.H. Krammer. 1989. Monoclonal antibody-mediated tumor regression by induction of apoptosis. Science. 245:301–305.

34. Brenner, D., A. Golks, F. Kiefer, P.H. Krammer, and R. Arnold. 2005. Activation or suppression of NFkappaB by HPK1 determines sensitivity to activation-induced cell death. EMBO J. 24:4279–4290.

35. Laemmli, U.K. 1970. Cleavage of structural proteins during the assembly of the head of bacteriophage T4. Nature. 227:680–685.

36. Medema, J.P., C. Scaffidi, F.C. Kischkel, A. Shevchenko, M. Mann, P.H. Krammer, and M.E. Peter. 1997. FLICE is activated by association with the CD95 death-inducing signaling complex (DISC). EMBO J. 16:2794–2804.

37. Klas, C., K.M. Debatin, R.R. Jonker, and P.H. Krammer. 1993. Activation interferes with the APO-1 pathway in mature human T cells. Int. Immunol. 5:625–630.