Human Procaspsase-1 Variants with Decreased Enzymatic Activity Are Associated with Febrile Episodes and May Contribute to Inflammation via RIP2 and NF-κB Signaling

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Human Procaspase-1 Variants with Decreased Enzymatic Activity Are Associated with Febrile Episodes and May Contribute to Inflammation via RIP2 and NF-κB Signaling

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The proinflammatory enzyme caspase-1 plays an important role in the innate immune system and is involved in a variety of inflammatory conditions. Rare naturally occurring human variants of the caspase-1 gene (CASP1) lead to different protein expression and structure and to decreased or absent enzymatic activity. Paradoxically, a significant number of patients with such variants suffer from febrile episodes despite decreased IL-1β production and secretion. In this study, we investigate how variant (pro)caspase-1 can possibly contribute to inflammation. In a transfaction model, such variant procaspase-1 binds receptor interacting protein kinase 2 (RIP2) via Caspase activation and recruitment domain (CARD)/CARD interaction and thereby activates NF-κB, whereas wild-type procaspase-1 reduces intracellular RIP2 levels by enzymatic cleavage and release into the supernatant. We approach the protein interactions by coimmunoprecipitation and confocal microscopy and show that NF-κB activation is inhibited by anti-RIP2-short hairpin RNA and by the expression of a RIP2 CARD-only protein. In conclusion, variant procaspase-1 binds RIP2 and thereby activates NF-κB. This pathway could possibly contribute to proinflammatory signaling. The Journal of Immunology, 2014, 192: 000–000.

The proinflammatory enzyme caspase-1 plays an important role in the innate immune system and is involved in a variety of inflammatory conditions such as periodic fevers or gout. Normally, inflammasome activation results in procaspase-1 auto-processing to generate the active form. This heterotetrameric enzyme containing two p10 and two p20 subunits is able to process the proforms of IL-1β and IL-18 into their biologically active mature forms (1–8). These cytokines subsequently participate in an appropriate response of the immune system to tissue damage or pathogen invasion (9) but also can play a role in autoinflammatory conditions (10).

Recently, we reported seven rare naturally occurring genetic variants of human procaspase-1, which exhibited altered protein structure and showed decreased or absent autoprocessing and enzymatic activity (11). However, such procaspase-1 variants with decreased activity were detected in patients with autoinflammatory disease. Therefore, we discussed possible disease modifying effects of the (pro)caspase-1 variants because they may have attenuated an otherwise more severe disorder (11). In addition, Lamkanfi et al. (12) showed that procaspase-1 with a defective active center can bind receptor interacting protein kinase 2 (RIP2) and thereby activate NF-κB. This NF is known as a key player in orchestrating inflammation (13). Therefore, we explored whether the naturally occurring procaspase-1 variants also can have a proinflammatory effect.
from Prof. J. Tschopp (University of Lausanne, Lausanne, Switzerland). The luciferase reporter plasmid pBxIVlacZ was a gift from Prof. G. Nolan (University of Michigan, Ann Arbor, MI). All plasmids for procaspase-1 variants have been described in Ref. 11. RIP2-wild-type (wt) was created by site-directed mutagenesis of the VSV-tagged RIP2-kinase-negative plasmid using the QuikChange II XL Site-Directed Mutagenesis Kit from Agilent Technologies (Santa Clara, CA), according to the manufacturer’s protocol. The CrmA plasmid was a gift from Dr. J. Bugert (Cardiff University, Cardiff, U.K.).

**Transfection protocol**

For transfection, HEK 293T cells were seeded in 6-well plates, transfected 24 h later with the respective plasmids with 1.75 μg/ml polyethylenimine, and evaluated 24 h later.

**NF-κB luciferase reporter assay**

NF-κB activity was determined using the Luciferase Assay System from Promega (Manheim, Germany), according to the manufacturer’s protocol. The activity of the luciferase was measured sequentially on a Mithras LB940 (Berthold, Bad Wildbad, Germany).

**Knockdown of RIP2 in HEK 293T cells**

HEK 293T cells were transfected with anti-RIP2 short hairpin RNA (shRNA) lentiviral particles (sc-37389-V) or control shRNA lentiviral particles (sc-108080); both from Santa Cruz Biotechnology using 3 μg/ml Polybrene, according to the manufacturer’s protocol. Cells were cultured in IMDM 10% FCS and selected with 4 μg/ml puromycin. Protein knockdown was analyzed by Western blotting using anti-RIP2 PX093 from Cell Sciences. Loaded protein was normalized to total protein concentration determined with DC Protein Assay from Bio-Rad (Munich, Germany).

**Immunofluorescence staining**

Human peripheral blood monocytes were isolated from PBMCs using CD14-magnetic beads from Miltenyi Biotec (Bergisch Gladbach, Germany) and differentiated to macrophages by incubation in M-CSF (50 ng/ml) for 7 d. The cells were then incubated with or without crude LPS (10 ng/ml) for 4 h. The cells were fixed in 4% paraformaldehyde for 10 min at 4˚C, permeabilized in PBS containing 0.04% saponin and 1% BSA for 1 h. Thereafter, the cells were stained with the appropriate primary Ab for 2 h (anti-RIP2, 8427, and anti–procaspase-1 A-19), followed by staining with the secondary Ab for 45 min. Cover slips were mounted on glass slides in Vectashield mounting medium from Vector Laboratories (Burlingame, CA). Cells were imaged using a Zeiss LSM 510 confocal microscope from Carl Zeiss (Jena, Germany) or a Leica TCS SP5 from Leica Microsystems (Wetzlar, Germany) with a ×40 or a ×63 1.4NA objective lens. An argon laser was used for excitation of Alexafluor 488 at wavelengths of 488 nm, whereas a helium-neon laser (543 nm) was used for excitation of Alexafluor 568.

Quantification of colocalization was performed with Fiji (14).

**Lentiviral transduction of THP-1 cells**

Expression of endogenous (pro)caspase-1 was knocked down using the shRNA expression vector pKO.1 (Addgene, Cambridge, MA). The following target sequences were used: CASP1-3’untranslated region, “AAGAGAT-CCTTCTGTAAGAGT”; and control, “AAGACCTCTTGTTAAGAGGT.” The knockdown was confirmed by Western blotting. An internal ribosome entry site (IRES) cassette was annexed to the Flag-tagged cDNA of wt or variant C285A-procaspase-1, and the constructs were cloned into the lentiviral transfer vector pRRL.SIN.cPPT.SFFV.GFP.WPRE [provided by Prof. C. Baum, Hannover Medical School, Institute of Experimental Hematology, Hannover, Germany (15)] with the GFP downstream of the IRES. To produce lentiviral vector particles, HEK 293T cells were transfected with lentiviral transfer plasmids in combination with the plasmids pSPAX2 and pVSVg. Next, the cells were harvested and infected with the supernatant 48 h after transduction. The infected cell media was concentrated by ultracentrifugation at 100,000 g for 2 h at 4˚C and resuspended in growth media (M199-10% FBS/P/S for the last 24 h). To study the release of RIP2, HEK 293T cells were transfected with plasmids coding for RIP2 and either wt procaspase-1 or the enzymatically inactive variant C285A. Proteins in the supernatant were precipitated with Polybrene, according to the manufacturer’s protocol. The precipitated proteins were analyzed by Western blotting using anti–procaspase-1 A-19. For immunoprecipitation, lysates were incubated with protein G-agarose from Santa Cruz Biotechnology overnight at 4˚C with rotation. Precipitates were analyzed by SDS-PAGE and Western blotting. All experiments were repeated at least three times.

**RIP2 cleavage experiments**

For RIP2 cleavage experiments, zVAD-FMK (50 μM) from Calbiochem (catalog number 219007; Merck Millipore, Billerica, MA) was added to the cell media 3 h prior harvesting. CrmA was supplied by cotransfection of a CrmA plasmid. RIP2 protein was detected by Western blot analysis using the anti-RIP2 Ab SP6266P.

In addition, the tandem fusion construct mCherry-RIP2-EGFP was introduced into the vector p6NST51. p6NST51 is a variant of p6NST50 (16) having the SFFV U3-driven IRES EGFP-Zeo expression cassette replaced by a IRES-Zeo resistance cassette. mCherry and EGFP genes have been purchased from BD Clontech (Mountain View, CA). RIP2 protein levels were determined by Western blot using anti-RIP2 Ab FPX093 (Cell Sciences) for the n-terminal RIP2 and anti-RIP2 Ab D10B11 (Cell Signaling Technology) for C-terminal RIP2.

**RIP2 release experiment**

To study the release of RIP2, HEK 293T cells were transfected with plasmids coding for RIP2 and either wt procaspase-1 or the enzymatically inactive variant C285A. Proteins in the supernatant were precipitated with methanol/chloroform. The cells had been cultured in media including insulin-transferrin-selenium-ethanolamine (Life Technologies) instead of FBS for the last 24 h.

**Statistics**

Significance of the NF-κB assay results and of Table I was determined by the paired Wilcoxon rank test (Holm adjusted) and Fisher’s exact test, respectively.

**Results**

**Procaspase-1 variants are associated with febrile episodes**

Recently, we characterized structure and function of different genetic variants of procaspase-1 (p.R221C, p.R240Q, p.N263S, p.L265S, p.T267I, p.K319R, and p.A329T) found in a healthy control population and in patients with recurrent febrile episodes (11). These procaspase-1 variants did not segregate with disease within families. Almost all relatives of patients with and without the variants were immunologically healthy. This excludes the variants from causing a monogenic disorder. However, we observed a statistically significant difference for the procaspase-1 variants to occur more often in patients than in healthy donors (Table I). Ethnicity is probably not a confounding factor because results are still significant within each group of Arabs, Turks, and whites. The rarest variants (R221C/wt, L265S/wt, T267I/wt, R240Q/240Q, and R240Q/K319R) occur in patients only.

Taken together, these results support the assumption that at least some of the variants are disease promoting cofactors.

**Procaspase-1 variants activate NF-κB**

Next, we analyzed a possible mechanism that could explain how the procaspase-1 variants contributed to proinflammatory signaling. For this purpose, we transfected HEK 293T cells with plasmids coding for wt and variant procaspase-1. The artificial variant C285A with a defective active center was included as control. The release of active NF-κB from the intrinsic NF-κB/IKB complex was measured by a luciferase reporter assay. The variant R221C was only weakly expressed at the protein level (11) and therefore

**Commonunoprecipitation using HEK 293T cells**

Cells were transfected in 6-well plates with RIP2-DSRed (800 ng) and/or variants of procaspase-1-Flag (200 ng) plasmids. After 24 h, cells were lysed with lysis buffer containing 20 mM Tris/HCl (pH 7.4), 150 mM NaCl, 10 mM EDTA, 0.2% Nonidet P-40, 1 mM PMSEF, and complete EDTA-free protease inhibitor (Roche Applied Science, Manheim, Germany). Lysates were clarified and analyzed by SDS-PAGE and Western blotting using anti-DSRed or anti–procaspase-1 A-19. For immunoprecipitation, lysates were incubated with anti-DSRed or anti–procaspase-1 A-19 for 2 h at 4˚C with rotation. Afterward, the immune complexes were incubated with protein G-agarose from Santa Cruz Biotechnology overnight at 4˚C with rotation. Precipitates were analyzed by SDS-PAGE and Western blotting. All experiments were repeated at least three times.

**Commonunoprecipitation using THP-1 cells**

THP-1 cells were differentiated with PMA (10 ng/ml) for 24 h and stimulated with 1 μg/ml crude LPS (L5293; purchased from Sigma-Aldrich). After incubation periods as indicated, cells were lysed with lysis buffer as described above, using anti-RIP2 N-19 for RIP2 precipitation. Densitometric analysis was performed using ImageJ (http://imagej.nih.gov/ij/).
not analyzed in this study. The other variants (N263S, K319R, R240Q, L265S, and C285A) and wt procaspase-1 were equally expressed after transfection (Fig. 1A). Furthermore, protein levels of endogenous RIP2 were also similar (Fig. 1A). The caspase-1 variants induced an increase in NF-κB activity that was significant in R240Q, L265S, and C285A (Fig. 1B). This increase roughly correlated inversely with enzymatic activity. Enhanced NF-κB activity was not accompanied by a increase or decrease of cell death rates of the transfected HEK 293T cells as measured by caspase-3 activation or ethidiumbromide staining of the transfected cells (Supplemental Fig. 1A, 1B). Hence, altered cell death rates do not explain differences in NF-κB activity.

The increase in NF-κB activity induced by variant procaspase-1 depends on RIP2

To investigate the participation of RIP2 in the observed enhanced NF-κB activation, we performed a cotransfection experiment with a plasmid coding for the CARD domain of RIP2 in HEK 293T cells. The coexpression of this plasmid alone abolished the pro-caspase-1–mediated NF-κB activation probably by blocking the

Table I. Procaspase-1 variants are associated with febrile episodes within different ethnic groups

<table>
<thead>
<tr>
<th>Ethnic Group</th>
<th>Patients: Variant Alleles/ Total Alleles</th>
<th>Control: Variant Alleles/ Total Alleles</th>
<th>Significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Whites</td>
<td>22/800</td>
<td>1/214</td>
<td>p = 0.03</td>
</tr>
<tr>
<td>Turks</td>
<td>7/102</td>
<td>2/200</td>
<td>p = 0.008</td>
</tr>
<tr>
<td>Arabs</td>
<td>3/12</td>
<td>4/200</td>
<td>p = 0.004</td>
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</tbody>
</table>


*Patients suffer from recurrent febrile episodes.

**Healthy blood donors.

FIGURE 1. NF-κB activation by procaspase-1 variants is mediated by RIP2. (A) Similarity of the expression levels of endogenous RIP2 as well as the different overexpressed procaspase-1 variants in HEK 293T was illustrated by Western blot using anti-RIP2 Ab D10B11 (C-terminal; Cell Signaling Technology) and anti–caspase-1 Ab A19 (Card-Domain; Santa Cruz Biotechnology). (B) Fold increase of intrinsic NF-κB activation in HEK 293T cells as measured by a luciferase reporter after transfection with plasmids coding for procaspase-1 variants as compared with transfection with empty plasmid (−, set to 1), n = 9. (C) Similar to (B), but cotransfected with a plasmid coding for the CARD domain of RIP2 where indicated; ordinate: arbitrary fluorescence units of luciferase substrate, n = 3. (D) Similar to (B), but transfection with plasmids coding for procaspase-1 variants after transduction of cells with anti-RIP2 shRNA (a-RIP2) or with irrelevant control shRNA (ctrl), n = 10. Inset confirms knockdown of RIP2 by a Western blot analysis with anti-RIP2 Ab PX093 (N-terminal; Cell Sciences).
interaction of the CARD domain of procaspase-1 with the CARD domain of complete RIP2 (Fig. 1C).

Furthermore, we knocked down RIP2 in HEK 293T cells with anti–RIP2-shRNA. Strongly decreased protein levels of RIP2 were confirmed (inset in Fig. 1D). Knockdown of RIP2 led to a significant reduction of NF-κB activation by the variants L265S and C285A compared with control cells (Fig. 1D). Possible shRNA off-target effects were excluded using anti-RIP2-shRNA constructs targeting two additional coding sequences of RIP2 (Supplemental Fig. 2).

Taken together, these data show that the procaspase-1 variants activate NF-κB via RIP2 in a transfection model.

Interaction and colocalization of procaspase-1 and RIP2 in THP-1 cells and human macrophages

In a next step, we examined whether endogenous procaspase-1 interacts with endogenous RIP2 in cells that belong to the immune system. Therefore, native THP-1 cells, differentiated with PMA and stimulated with crude LPS (cLPS) for different time intervals, were lysed, and endogenous RIP2 was immunoprecipitated. We found that endogenous procaspase-1 coprecipitated in a time-dependent manner (Fig. 2A). The strongest RIP2–procaspase-1 interaction occurred after 60–80 min and decreased after ∼3 h.

Next, we localized RIP2 and procaspase-1 in human monocyte-derived macrophages using Ab staining and confocal microscopy. After stimulation with cLPS, RIP2 and procaspase-1 colocalized in a cytoplasmic compartment adjacent to the nucleus of the macrophages. This happened in a time dependent manner and in accordance with the communoprecipitation (co-IP) experiment (Fig. 2A). The highest rate of cells showing this colocalization was found 60–80 min after the beginning of cLPS stimulation (Fig. 2B, 2C). Medium exchange alone did not induce colocalization of RIP2 and procaspase-1 (Supplemental Fig. 3). The colocalization diminished over ∼3 h and then reached the level of nonstimulated cells. The localization of procaspase-1 and RIP2 in the same area was further ensured by determining Pearson’s correlation coefficient [which will be close to 1 if green and red channel distributions are linked (17)] and Manders’ overlap coefficient (Fig. 3A). In accordance with the cell staining, fluorescence intensity profiles of both colors correlated in a corresponding histogram (Fig. 3B).

Analysis of interaction of procaspase-1 variants with RIP2 by Co-IP

To investigate the interaction of RIP2 with an inactive procaspase-1 variant, we established THP-1 cells that had been transduced with shRNA targeting a sequence of the 3′-untranslated region of wt procaspase-1 mRNA to knock down endogenous wt procaspase-1. The reconstitution with either wt procaspase-1 or the enzymatically inactive variant L265S was realized by a second transduction. Western blot analysis showed similar amounts of introduced procaspases compared with endogenous procaspase-1 in control cells (Fig. 4A, lysate, middle row).

Prior to IP, the cells were differentiated with PMA and stimulated with cLPS. Endogenous RIP2 levels were constant in the cell lysates (Fig. 4A, lysate, upper row) and in the immunoprecipitated fraction (Fig. 4A, IP: aRIP2, upper row). The IP of intrinsic RIP2 led to coprecipitation of reconstituted procaspase-1 (Fig. 4A, IP: a-RIP2, bottom row). However, interaction of the enzymatically inactive variant L265S with RIP2 was increased with wt procaspase-1 (Fig. 4A, IP: a-RIP2, bottom row). Although the enzymatically inactive variant was expressed 1.3-fold stronger than reconstituted wt procaspase-1m the lysate, the precipitation of the L265S variant with RIP2 was 2.3-fold higher as determined densitometrically.

In parallel, we cotransfected HEK 293T cells with plasmids coding for VSV-tagged RIP2 and either wt procaspase-1 or one of

![FIGURE 2.](http://www.jimmunol.org/) Co-precipitation and colocalization of endogenous procaspase-1 with RIP2 in human macrophages. (A) THP-1 cells were differentiated to macrophages with PMA (10 ng/ml, 24 h) and stimulated with LPS (1 μg/ml) for different time intervals. Lysate: constant expression of RIP2 and procaspase-1. Co-IP of endogenous procaspase-1 with endogenous RIP2 shows a time course of interaction. (B) Quantification of colocalization of procaspase-1 and RIP2 in monocyte-derived macrophages. A total of 1000 cells in three independent experiments has been analyzed for colocalization of procaspase-1 and RIP2. Colocalization-positive cells as shown in (C) have been counted after different time intervals of LPS (10 ng/ml) stimulation and are shown as percentage of total cells. Note the similarity in the time course of Co-IP in (A) and of colocalization in (B). (C) Ab staining and confocal microscopy of procaspase-1 and RIP2 in human monocyte-derived macrophages. Original magnification ×63. After stimulation with LPS, RIP2 and procaspase-1 colocalize perinuclear in the cytoplasm of the macrophages (white arrows). RIP2 is shown in the red, procaspase-1 in the green, and nuclei in the blue (DAPI) channel. Left panel, nonstimulated cells, right panel, cells after 80-min incubation with LPS. Colocalization was also seen with Abs binding to epitopes of procaspase-1 not located in the CARD domain (data not shown).

the enzymatically inactive variants L265S and C285A to gain higher levels of these proteins within cells. Again, IP of VSV-RIP2 led to coprecipitation of procaspase-1 and vice versa (Fig. 4B).
However, we noticed that cotransfection with a plasmid coding for wt procaspase-1 yielded much fainter VSV-RIP2 bands compared with cells cotransfected with plasmids coding for L265S or C285A variants (Fig. 4B, lysate, upper row). The differences of the VSV-RIP2 protein levels became even stronger in the co-IP (Fig. 4b, a-procaspase-1 IP). Similar to the anti-RIP2 IP with the transduced THP-1 cells, anti–VSV-RIP2 IP in transfected HEK 293T cells yielded much stronger bands of enzymatically inactive procaspase-1 variants compared with wt procaspase-1 (Fig. 4B, a-VSV IP).

These data indicate that interaction of enzymatically inactive procaspase-1 variants with RIP2 was enhanced.

wt but not variant procaspase-1 decreased intracellular RIP2 levels by enzymatic cleavage and by release of RIP2 into the supernatant

The previous experiments suggested decreased VSV-RIP2 protein levels in cells cotransfected with a plasmid coding for wt procaspase-1. Next, we investigated this phenomenon. First, plasmids coding for different procaspase-1 variants were transfected together with a plasmid coding for VSV-RIP2 into HEK 293T cells. In Western blot analysis, wt and enzymatically residual active procaspase-1 variants (e.g., N263S) yielded indeed fainter bands for VSV-RIP2 than inactive procaspase-1 forms (Fig. 5A). Moreover, an 8-kDa C-terminal cleavage product of RIP2 could be found when active procaspase-1 had been cotransfected (Fig. 5B).

Second, cleavage experiments with the tandem fusion protein mCherry-RIP2-EGFP (cartoon in Fig. 5B) revealed another RIP2 cleavage product at the N terminus fused to mCherry (80 kDa; Fig. 5B, first row). The C-terminal cleavage product fused to GFP (30 kDa) could be confirmed (Fig. 5B, second row).

Third, two different kinds of caspase-inhibitors, zVAD-CMK and CrmA, prevented the wt procaspase-1–mediated reduction of VSV-RIP2 protein level in the lysate (Fig. 5C).

Finally, we tested the hypothesis that coexpression of wt procaspase-1 with VSV-RIP2 induces the release of VSV-RIP2 into the supernatant. As shown in Fig. 5D, VSV-RIP2 could be detected in the supernatant of cells, which had been cotransfected with a plasmid coding for wt procaspase-1 in contrast to cells with the inactive variant C285A or without procaspase-1 (Fig. 5D, supernatant). This release of VSV-RIP2 correlated with a reduction of intracellular VSV-RIP2 protein level (Fig. 5D, lysate). Activation
of wt procaspase-1 was confirmed by the detection of p20 in the lysate, independent of VSV-RIP2 cotransfection. The p20 subunit was not detected in cells expressing the inactive variant C285A (Fig. 5D, lysate). However, p20 was only released into the supernatant of cells, which had been cotransfected with a plasmid coding for wt RIP2 and procaspase-1 variants, and addition of the caspase inhibitors zVAD and CrmA (by cotransfection). The multiple bands in the lysate were generated by cutting RIP2 and also the annexed dyes (anti-RIP2 Ab PX093, anti-GFP Ab, data not shown, in this study stained with anti-GAPDH Ab, data not shown, in this study stained with anti-GAPDH Ab). The tandem protein mCherry-RIP2-EGFP is cut by wt procaspase-1 yielding a C-terminal (C-t.) cleavage product of ∼30 kDa from RIP2 that is fused to GFP (the band in the second row also can be stained with an anti-GFP Ab, data not shown, in this study stained with anti-RIP2 Ab D10B11, C-terminal). The multiple bands in the first row are also generated by cutting RIP2 and also the annexed dyes (anti-RIP2 Ab PX093, N-terminal). Similar Western blot as in A after cotransfection with plasmids coding for wt RIP2 and wt procaspase-1, and addition of the caspase inhibitors zVAD or CrmA (by cotransfection).

FIGURE 5. RIP2 gets cleaved and released from the cells dependent on the enzymatic activity of caspase-1. (A) Western blot with anti-RIP2 Ab (SP6266P, C-terminal) after cotransfection of HEK 293T cells with plasmids coding for wt RIP2 and procaspase-1 variants shows a RIP2 cleavage product depending on enzymatic activity of caspase-1. (B) Confirmation of N-terminal (N-t.) cleavage product of RIP2. The tandem protein mCherry-RIP2-EGFP is cut by wt procaspase-1 yielding a C-terminal (C-t.) cleavage product of ∼30 kDa from RIP2 that is fused to GFP (the band in the second row also can be stained with an anti-GFP Ab, data not shown, in this study stained with anti-RIP2 Ab D10B11, C-terminal). The multiple bands in the first row are generated by cutting RIP2 and also the annexed dyes (anti-RIP2 Ab PX093, N-terminal). (C) Similar Western blot as in A after cotransfection with plasmids coding for wt RIP2 and wt procaspase-1, and addition of the caspase inhibitors zVAD or CrmA (by cotransfection). (D) Western blot of lysate and precipitated supernatant of HEK 293T cells transfected with plasmids coding for VSV-RIP2 and procaspase-1 wt or the enzymatically inactive variant C285A. The protein levels had been determined using the following Abs: anti-RIP2 Ab PX093 (N-terminal; Cell Sciences), anti–caspase-1–cleaved p20 Ab h297 (Santa Cruz Biotechnology) for cleaved caspase-1 subunit p20 (p20), anti–caspase-1 cleaved p20 Ab C285A, anti–GAPDH Ab H86045M (Meridian Life Science), and anti–caspase-1 cleaved p20 Ab C285A (Santa Cruz Biotechnology) for procaspase-1 p45 (p45).

Discussion

Recently, we have described human hypomorphic genetic variants of procaspase-1 (11). Phagocytes and especially monocytes from individuals with such variants show a trend to produce decreased amounts of active IL-1β (11). Therefore, such individuals should be less prone to develop autoinflammation than others. Unexpectedly, we find a significant correlation between enzymatic less or inactive procaspase-1 variants and the development of autoinflammatory disorders (Table I). In an attempt to solve this paradox, we refer to the suggestion of Lamkanfi et al. (12) and investigate the ability of the naturally occurring human procaspase-1 variants to activate NF-κB via RIP2 interaction.

In a transfection model with endogenous RIP2, we find indeed a robust activation of intrinsic NF-κB by naturally occurring hypomorphic procaspase-1 variants (Fig. 1B). Interestingly, this activation is roughly inversely proportional to the residual enzymatic activity of the variants. It depends on RIP2 and is mediated by CARD/CARD interaction between procaspase-1 and RIP2 (Fig. 1C, 1D). In accordance with this interaction, endogenous RIP2 and procaspase-1 coimmunoprecipitate (Fig. 2A). Moreover, the endogenous proteins colocalize after activation of monocyte-derived macrophages (Figs. 2C, 3) and therefore putatively interact and take part in the same molecular pathways. Using transduction of THP-1 cells with procaspase-1 variants by lentiviral vectors with limited copy numbers per cell, we avoid strong protein overexpression and approximate a physiologic situation (Fig. 4A). As indicated by coimmunoprecipitation, interaction of the enzymatically inactive procaspase-1 variant L265S with endogenous RIP2 is stronger than the respective interaction with wt procaspase-1. Co-IP of procaspase-1 and VSV-RIP2 coexpressed in transfected HEK 293T cells strongly supports the data obtained in THP-1 cells. Enzymatically inactive procaspase-1 variants interact much stronger with RIP2 than wt procaspase-1 (Fig. 4B).

The activation of NF-κB by enzymatic inactive procaspase-1 described in our study is in accordance with the data from Lamkanfi et al. (12). However, in contrast to their study, we avoid wt procaspase-1–mediated cell death (Supplemental Fig. 1A) by reducing caspase-1 expression levels. Moreover, we show equal protein

FIGURE 6. Model for proinflammatory effects induced by procaspase-1 variants with reduced enzymatic activity. The procaspase-1 variants exhibit reduced or absent enzymatic activity because of the different missense mutations, which are substituted by asterisks (from the left to the right: p.R221C, p.R240Q, p.N263S, p.L265S, p.T267I, p.K319R, and p.A329T). Although procaspase-1 autoprocessing and IL-1β activation is decreased by such procaspase-1 variants (bottom), the NF-κB activation is increased (top) as a result of the longer and/or stronger interaction between procaspase-1 and RIP2. The enhanced NF-κB activity may contribute to the transcription of proinflammatory proteins and thereby aggravate or unmask inflammatory disorders.
levels of wt procaspase-1 and all variants analyzed for NF-κB activation (Fig. 1A). Thereby, we are able to compare the NF-κB activating potential of different procaspase-1 variants with wt procaspase-1. We show an increased NF-κB activation mediated by variant procaspase-1. The underlying mechanism may be similar to NF-κB activation after binding of RIP2 to nucleotide-binding oligomerization domain-containing protein 2 (18). Hypothetically, the procaspase-1 variants with residual or no enzymatic activity induce longer NF-κB activation compared with wt procaspase-1 mediated by a longer and/or stronger interaction with RIP2. NF-κB can subsequently bind to regulatory sequences of genes that encode proinflammatory proteins and thereby upregulate their expression (19). 

In contrast to enzymatically inactive procaspase-1, wt procaspase-1, and procaspase-1 variants with residual activity (e.g., N263S) reduce intracellular levels of overexpressed RIP2 (Fig. 5A). This effect as well as the faint VSV-RIP2 bands in the immunoprecipitation experiments (Fig. 4B) can be explained by RIP2 cleavage and release from cells induced by wt procaspase-1 (Fig. 5). The activation of caspase-1 and the simultaneous release of the p20 subunit by cotransfection of plasmids coding for wt procaspase-1 with RIP2 indeed has already been described (19). In this study, we additionally show the release of the overexpressed RIP2 from the cells. To our knowledge, cleavage of RIP2 by endogenous proteases has not been published to date.

So far, reduced enzymatic activity of caspase-1 realized by inhibition or knockout in mice is mainly thought to decrease inflammation (20–23). Considering the data of Kayagaki et al. (24), most cells induced by wt procaspase-1 (Fig. 5). The activation of caspase-1 as well as the faint VSV-RIP2 bands in the immunoprecipitation experiments (Fig. 5A) can be attributed to RIP2 cleavage and release from cells induced by wt procaspase-1 with RIP2 indeed has already been described (19). In this study, we additionally show the release of the overexpressed RIP2 from the cells. To our knowledge, cleavage of RIP2 by endogenous proteases has not been published to date.

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Suppl. Fig. 1: Procaspase-1 overexpression with endogenous protein levels of RIP2 does not increase cell death.

A: Caspase-3 activity, indicating apoptosis, was measured in HEK 293T cells transiently transfected with plasmids coding for procaspase-1 variants. For comparison: e.p. transfection with empty plasmid, J.c. non-induced Jurkat cells, J.m. Jurkat cells incubated with agonistic proapoptotic mAb CH11. Caspase-3 activity was determined using the Enz Check Caspase-3 Assay Kit #1 (Molecular Probes, Leiden, The Netherlands). B: Ethidium bromide stained nuclei, indicating membrane damage, were determined in HEK 293T cells transiently transfected with a plasmid coding for wildtype procaspase-1 (wt) or the enzymatically inactive variant C285A; for comparison: e.p. transfection with empty plasmid, pos. ctrl.: untransfected cells permeabilized with Triton X-100. The ratio of cells with ethidium bromide stained nuclei to cells with generally stained nuclei (Hoechst) represent the ratio of cells with a defective cell membrane.
Suppl. Fig. 2: NF-κB activation by procaspase-1 variants is decreased by two additional anti-RIP2 shRNAs.

**A:** Fold increase of intrinsic NF-κB in HEK 293T cells as measured by a luciferase reporter after transduction of cells with anti-RIP2 shRNA (a-RIP2 #2 or a-RIP2 #3) or with irrelevant control shRNA (ctrl) and transient transfection with a plasmid coding for wildtype procaspase-1 (wt) or the enzymatically inactive variant C285A as compared to transfection with empty plasmid (-; set to 1), n≥9. Insert confirms knock down of RIP2 by a Western blot with anti-RIP antibody PX093 (Cell Sciences, N-terminal). The target shRNAs were obtained from TRC shRNA library and the following sequences were used: anti-RIP2 shRNA #2: “GCACCATTCTGGATCTCAAA”; anti-RIP2 shRNA #3: “GCACAATATGACTCCTCCTTT”; control shRNA, “TACAACAGCCACAACGTCTAT”.
Suppl. Fig. 3: Co-localisation of Procaspase-1 with RIP2 in primary human macrophages. A: Antibody staining and confocal microscopy of primary human macrophages. Human monocytes were isolated from peripheral blood by CD14 selection and subsequently differentiated with M-CSF to macrophages. After 40 to 80 min of stimulation with MDP + upLPS, RIP2 and procaspase-1 co-localized perinuclear in the cytoplasm of the macrophages (white arrows). RIP2 is shown in the red channel, procaspase-1 in green and
Dapi in blue. Cells with medium exchange only are shown in the left panel and cells after MDP + upLPS in the right panel. **B:** Quantification of co-localisation. A total of 100 cells per sample has been analysed for co-localisation of procaspase-1 and RIP2. For co-localisation, positive cells have been counted and shown as percent compared to total cells. Cells receiving medium exchange only at given time points did not show co-localisation (or in a very low level). The highest number of cells showing co-localisation after treatment with MDP + LPS was determined 120 minutes after the start of stimulation, and diminished thereafter. upLPS - ultrapure LPS, MDP – muramyldipeptide.