Eczematous dermatitis: T cell and keratinocyte apoptosis plays a key pathogenetic role.

Axel Trautmann¹, Mübeccel Akdis², Daniela Kleemann², Frank Altznauer³, Hans-Uwe Simon¹, Thomas Graeves⁴, Michaela Noll³, Eva-B. Bröcker⁴, Kurt Blaser¹ and Cezmi A Akdis¹

¹Swiss Institute of Allergy and Asthma Research (SIAF), Davos, Switzerland
²High-altitude Clinic of Zürich, Davos-Clavadel, Switzerland
³Fraunhofer Institute for Interfacial Engineering and Biotechnology, Stuttgart, Germany
⁴Department of Dermatology, University of Würzburg, Würzburg, Germany

*Corresponding Author: Axel Trautmann, Swiss Institute of Allergy and Asthma Research (SIAF), Obere Strasse 22, CH-7270 Davos, Switzerland. Phone: 41-81-4100848; Fax: 41-81-4100840; E-mail: hc.hzinu.fais@tutarra.ch

Cite this article: Eczematous dermatitis: T cell and keratinocyte apoptosis plays a key pathogenetic role. Sci J of Der and Ven. 2018; 1(1): 001-007

Submitted: 03 February 2018; Approved: 24 February 2018; Published: 25 February 2018

ABSTRACT

Clinical and histologic similarities between various eczematous disorders point to a common effector pathway. We demonstrate here that activated T cells, infiltrating the skin in atopic dermatitis (AD) and allergic contact dermatitis (ACD), induce keratinocyte (KC) apoptosis. KCs normally express low levels of Fas receptor (FasR) that can be substantially enhanced by the presence of IFN-gamma. KCs are rendered susceptible to apoptosis by IFN-gamma when FasR numbers reach a threshold of approximately 40,000 per KC. Subsequently, KCs undergo apoptosis induced by anti-FasR mAb's, soluble Fas ligand, supernatants from activated T cells, or direct contact between T cells and KCs. Apoptotic KCs show typical DNA fragmentation and membrane phosphatidylserine expression. KC apoptosis was demonstrated in situ in lesional skin affected by AD, ACD, and patch tests. Using numerous cytokines and anti-cytokine neutralizing antibodies, we have shown that cytokines and cell-surface molecules of the accumulated skin-infiltrating T cells affect KCs in situ. In this study, we demonstrated that KC apoptosis is a major mechanism in the pathogenesis of eczematous disorders. T cells infiltrating the skin upregulate Fas receptor (FasR) on KCs, render them susceptible to apoptosis by IFN-γ, and induce apoptosis by Fas ligand (FasL) expressed on the T-cell surface.

INTRODUCTION

Eczematous disorders account for a large proportion of all skin disease, and constitute a major health problem worldwide. Eczematous dermatitis is characterized histologically by pathologic changes in the epidermis and a predominantly mononuclear cell infiltrate. The clinical features include itching, redness, papules, vesicles, and scaling. Because T cells constitute a large population of cellular infiltrate in eczematous dermatitis, a dysregulated, cytokine-mediated response of the immune system appears to be an important pathogenetic factor: Keratinocytes (KC's) within eczematous lesions exhibit an unusual expression of MHC class II antigens (e.g., HLA-DR, ICAM-1 (CD54)), and IFN-γ inducible protein-10, all of which are strongly induced by IFN-γ (1, 2). ICAM-1 mediates strong adhesion between T cells and KCs (3). Thus it seems probable that cytokines and cell-surface molecules of the accumulated skin-infiltrating T cells affect KCs in situ.

Atopic dermatitis (AD) is a chronic skin disease manifested by eczematous skin lesions that affects more than 10% of children (4, 5). Lesional atopic skin clinically and histologically resembles allergic contact dermatitis (ACD), which may be mediated by predominant expression of the type 1 cytokine IFN-γ (6, 7). In AD, circulating allergen-specific memory effector T cells expressing cutaneous lymphocyte-associated antigen homing receptor have been demonstrated to be activated in vivo, to regulate IgE by an IL-13-dominated cytokine pattern, and to delay eosinophil apoptosis by IL-5 (8–10). Allergen-specific T cells from skin and peripheral blood have shown a profound dysregulation of IL-4 and IFN-γ secretion, with an overproduction of IL-4 and reduced IFN-γ secretion in AD (11, 12). However, studies focused on the intracellular cytokine production revealed a higher IFN-γ production than occurs in T cells from peripheral blood (13, 14). A sequential activation of type 2 and type 1 T cells in the pathogenesis of AD was therefore proposed (15–17). Recent results from a murine model of eczematous dermatitis suggests that both the type 2 cytokines IL-4 and IL-5, and the type 1 cytokine IFN-γ play important roles in the inflammation of the skin (18). Furthermore, injection of IFN-γ into the skin of human volunteers demonstrated that this cytokine can induce transient skin inflammation (19). Moreover, transgenic mice expressing IFN-γ in the epidermis have developed eczema spontaneously, and showed an increased contact hypersensitivity reaction (20).

METHODS

Subjects

Patients with acute AD, chronic AD, and acute ACD were included in the study. They were not receiving any systemic therapy for at least 2 weeks before biopsy and patch testing with house dust mite and nickel. AD patients had definite AD, diagnosed according to standard criteria (21). ACD for nickel was originally diagnosed by epicutaneous patch testing. The patch test reactions were 2+ according to the International Contact Dermatitis Research Group, showing erythema, induration, papules, and vesicles. Incisional skin biopsies were taken from lesional AD skin (acute AD, four biopsies; chronic AD, five biopsies), atopic patch tests (two biopsies), and ACD patch tests (four biopsies). Control skin was obtained from three healthy, nonatopic individuals. Informed consent was obtained from all subjects, and the study was approved by the Ethical Committee of Davos, Switzerland.

Reagents and Abs

Recombinant human IL-12 and IL-5 were purchased from Pharmingen (San Diego, California, USA). Recombinant human IL-13 was purchased from PeproTech Inc. (Rocky Hill, New Jersey, USA). Recombinant human IFN-γ, IL-2, IL-4, and TNF-α came from Novartis Pharmaceuticals (Basel, Switzerland). Ethidium bromide and PMA were purchased from Sigma Chemical Co. (St. Louis, Missouri, USA). Ionomycin was purchased from Calbiochem-Novabiochem Corp. (San Diego, California, USA). The mAb’s for flow cytometry were purchased from Beckman Coulter Intl. SA (Nyon, Switzerland). Immunotech (Marseilles, France), or Pharmingen. Anti-CD14, anti-CD16, anti-CD19, and anti-CD45RA magnetic microbeads for magnetic activated cell sorting (MACS) were from Miltenyi Biotec (Bergisch Gladbach, Germany). Anti-CD11a, anti-CD11b, anti-CD11c, anti-CD34, and anti-CD45RA magnetic microbeads for magnetic activated cell sorting (MACS) were from Miltenyi Biotec (Bergisch Gladbach, Germany).
The blocking anti–IFN-γ receptor–α chain (CD119) mAb was purchased from Genzyme Pharmaceutica~(Cambridge, Massachusetts, USA). Neutralizing anti–IL-4 mAbs (8B12 and 3H4) and IFN-γ mAb 45-15 were provided by Novartis Pharmaceutica~.

Anti–IL-13 mAb (JES8-5A2 and JES10-2P9) and the mutant IL-4 antagonist Y124D with IL-13- and IL-4-inhibitory activity were provided by DNA Research Institute (Palo Alto, California, USA). The recombinant human soluble Fasl and Fas-Fc protein (inhibiting the activity of FasL) were purchased from Alexis Corp. (San Diego, California, USA). The enzyme inhibitor (Z)-Val-Ala-dl-Asp-fluoromethylketone (Z-VAD-FMK) was from Bachem AG (Bubendorf, Switzerland).

**KC Culture**

Primary human KCs were obtained from punch biopsies from nonlesional, nonmedicated gluteal skin. The skin was split overnight in a PBS solution of sucrose and trypsin (0.1% sucrose, 0.25% trypsin, 1 mM EDTA; all reagents were from Sigma Chemical Co.) at 4°C. Epidermal sheets were removed from the dermis, and KC cell suspensions were cultured in a fully supplemented (5 μg/mL bovine insulin, 0.5 μg/mL hydrocortisone, 0.1 ng/mL human recombinant EGF, 30 μg/mL bovine pituitary extract, 100 μg/mL gentamicin, and 100 ng/mL amphotericin B), low calcium (0.15 mM Ca2+), serum-free KC growth medium (Clonetics Corp., San Diego, California, USA). Hydrocortisone and antibiotics were removed from the culture medium for the duration of experiments. All cell cultures were incubated at 37°C in a humidified atmosphere with 5% CO2. KCs that had undergone at least three passages were grown in either 6-well Transwell plates, 6-well plates, or 96-well flat-bottom plates (all from Corning-Costar Corp., Cambridge, United Kingdom) for different experiments. Single-cell suspensions of KCs were prepared by the addition of 0.025% trypsin and 0.01% EDTA for 5–15 minutes at 37°C, neutralized by adding 10% FCS (Sera-Lab Ltd., Sussex, United Kingdom).

Isolation of CD45RA+ and CD45RO+ T cells from peripheral blood and differentiation and characterization of CD4+ and CD8+ type 1 and type 2 T cells.

After 2–3 weeks of KC culture, PBMCs from the same patient were isolated by Ficoll (Biochrom KG, Berlin, Germany) density gradient centrifugation of peripheral venous blood. Cells were washed and resuspended in DMEM supplemented with 10 mM HEPES (both from Life Technologies AG, Basel, Switzerland), 5 mM EDTA, 2% FCS, 100 U/mL glutamine, and 100 μg/mL streptomycin (both from Life Technologies AG). CD45RA+ and CD45RO+ T cells were isolated with the MACS system according to the instructions of the manufacturer (Miltenyi Biotec). In brief, anti-CD4– and anti-CD19–depleted cells were incubated with MACS microbeads coated with anti-CD45RA and anti-CD16 mAb, and the T-cell fractions were recovered by sequential elution from the MACS column. CD4+ and CD8+ type 1 and type 2 T cells were generated from naive CD45RA+ T cells isolated from cord blood. The cells were incubated in the presence of either IL-12 (10 ng/mL) and neutralizing anti–IL-4 mAb (10 μg/mL) (for the generation of type 1 T cells) or IL-4 (25 ng/mL) and neutralizing anti–IL-12 mAb (10 μg/mL) (for the generation of type 2 T cells) in the presence of IL-2 (20 ng/mL) for 12 days. The T cells were stimulated with a combination of mAbs (0.5 μg/mL anti-CD2, 1.0 μg/mL anti-CD3, and 0.5 μg/mL anti-CD28). Supernatants were harvested after 3 days for cytokine measurements. Phenotypes of type 1 and type 2 T-cell subsets were verified for cytokine production by ELISA of culture supernatants and intracellular cytokine flow cytometry analysis.

**KC–T Cell Cocultures**

CD45R0+ T cells and differentiated type 1 and type 2 T cells were stimulated with a combination of anti-CD2, anti-CD3, and anti-CD28 mAbs, plate-bound anti-CD3 (10 μg/mL), or PMA (10 ng/mL) and ionomycin (0.5 μM) for 24 hours, and then washed twice before coculture with KCs. In some experiments, type 2 T cells were stimulated in the presence of IL-12 (10 ng/mL). For studying the effects of T cells on KCs, T-cell supernatants were collected, diluted 50%, and added to KCs in 96-well plates. For coculture experiments, KCs were cultured in 96-well plates in the presence or absence of purified T cells at different ratios (generally 104 T cells and 3 × 104 KCs, ratio 1:3). For further coculture experiments, KCs and T cells were cultured in 6-well Transwell plates (Corning-Costar Corp.). The Transwells consist of a lower and an upper compartment, which are separated by a 10-μm-thick polycarbonate membrane with 0.4-μm pores. KCs (3 × 105) were transferred into the lower culture well and T cells (105) were added to the upper compartment.

**Flow Cytometry Analysis**

IFN-γ receptor and IFN-γ-induced FasR (CD95) expression was determined on primary human KCs with anti–IFN-γ receptor and FasR mAb (ZB4), followed by phycoerythrin-conjugated (PE-conjugated) anti-mouse IgG. For phenotyping of effector T cell subsets, 5 × 105 cells were incubated at 4°C for 30 minutes, and then sequentially stained with anti-CD4+, anti-CD8– or CD45RO– PE. Stained cells were fixed in 2% paraformaldehyde. PE-conjugated mouse IgG1 was used as control. After washing, cells were analyzed on an EPICS XL-MCL flow cytometer (Beckman Coulter Intl. SA). The purity of the T-cell populations ranged from 90% to 96%. FasR counting on KC surfaces was done by indirect immune fluorescence staining with 50 μg/mL anti-FasR mAb (ZB4) followed by FITC-conjugated anti-mouse IgG (DAKO A/S, Glostrup, Denmark). Beads coated with known amounts of mouse mAb’s were used as standards (DAKO A/S) (22). The receptor molecules per cell were probably counted in lower numbers with this method than the classical Scatchard blot receptor determination would yield.

**Viability and Apoptosis Detection**

The KC viability after exposure to anti-FasR mAb, soluble Fasl, or after coculture with T cells was routinely evaluated by means of ethidium bromide (1 μM) uptake and flow cytometry. Membrane phosphatidylserine redistribution from the inner to the outer membrane leaflet takes place in apoptotic cells. Annexin V is a phosphatidylserine binding protein used to detect apoptotic cells. The technique was performed according to Vermes et al. (23). Briefly, cells were incubated with 1.0 μg/mL annexin V–FITC (R&D Systems Inc., Abingdon, United Kingdom) and 2.5 μg/mL propidium iodide (Sigma Chemical Co.) in calcium-containing binding buffer (HEPES with 0.25 mM CaCl2). DNA flow cytometry was performed according to Nicoletti et al. (24). In brief, cells were resuspended in hypotonic fluorochrome solution containing 50 μg/mL propidium iodide (Sigma Chemical Co.), 0.1% sodium citrate (Fluka Chemie AG, Buchs, Switzerland), and 0.1% (vol/vol) Triton-X 100 (Fluka Chemie AG), and then incubated at 4°C for 6 hours.

**Rnase Protection Assay**

Third-passage primary human KCs were treated with FasR mAb, recombinant human IFN-γ, and supernatant from stimulated CD45RO+ T cells diluted 50%. The KCs were lysed on the plate, and the RNA was isolated according to the RNaseasy protocol (QIAGEN AG, Basel, Switzerland), which includes DNase digestion. The RNase protection assay was performed according to the protocol of the manufacturer (PharMingen). Briefly, all RNA obtained from 3 × 106 cells (3–4 μg) was hybridized overnight to the 32P-labeled RNA probe (hApo3c; PharMingen). Single-stranded RNA and free probe were digested by RNase A and RNase T1. Protected RNA was collected and hybridized to a membrane filter. Inspection of the filter by autoradiography was performed according to the供应商的提示。The technique was performed according to Nicoletti et al. (24). In brief, cells were resuspended in hypotonic fluorochrome solution containing 50 μg/mL propidium iodide (Sigma Chemical Co.), 0.1% sodium citrate (Fluka Chemie AG, Buchs, Switzerland), and 0.1% (vol/vol) Triton-X 100 (Fluka Chemie AG), and then incubated at 4°C for 6 hours.

**Quantification of Cytokines**

IL-4, IL-5, IL-13, and IFN-γ were determined by sandwich ELISA as described (9, 10). The sensitivity of IL-4 ELISA was at or below 20 pg/mL (mAb and IL-4 standard were provided by C.H. Heusser, Novartis Pharmaceutica~). The detection limit of the IL-5 ELISA was 50 pg/mL (mAb and IL-5 standard were from PharMingen). The detection limit of IL-13 ELISA was 100 pg/mL (mAb and IL-13 standard were from PharMingen). The sensitivity of the...
IFN-γ ELISA was at or below 10 pg/mL (mAb and IFN-γ standard were gifts from S.S. Alkan, Novartis Pharmaceuticals). Soluble FasL was detected by a commercial ELISA kit (MBL Co., Nagoya, Japan) with a sensitivity of 0.1 ng/mL.

**Immunohistology**

The tissue samples were placed in Tissue-Tek OCT compound (Sakura Finetek Europe BV, Zoeterwoude, The Netherlands) and stored at −80°C. Four-micrometer cryostat sections were prepared on gelatin-coated slides (Merck AG, Dietikon, Switzerland). After air drying, sections were fixed in acetone for 10 minutes at 4°C. A three-step streptavidin-biotin complex peroxidase method (streptABCComplex-peroxidase; DAKO A/S) was used. Briefly, sections were incubated with the primary mAb (CD4, CD8, and CD45RO from DAKO A/S; FasL from Alexis Corp.; IFN-γ receptor from Genzyme Pharmaceuticals; and FasR, anti-perforin, and anti-granzyme B from Ancell Corp., Bayport, Minnesota, USA) at 4°C overnight. This was followed by incubation with biotin-conjugated rabbit anti-mouse IgG and preformed streptABCComplex-peroxidase (both from DAKO A/S) at room temperature for 1 hour. Incubation with the peroxidase-specific substrate 3-aminoo-9-ethylcarbazole (Sigma Chemical Co.) was used for visualization, with hematoxylin counterstaining. For control purposes, the primary mAb was replaced by an irrelevant isotype-matched mAb that consistently yielded negative results.

**Identification of Apoptosis by HOECHST Staining**

HOECHST staining was done according to Norris et al. (26) in KCs cultured in Transwell plates (Corning-Costar Corp.) and in lesional skin sections. After fixation with 4% paraformaldehyde and 200 mM dihydrogen phosphate (pH 7.0) overnight at 4°C, staining was performed with HOECHST 33342 dye (1 μg/mL; Sigma Chemical Co.) for 5 minutes. Stained sections were evaluated with an ultraviolet microscope (Axiovert 405M; Carl Zeiss AG, Feldbach, Switzerland).

**Cocultures of Normal Human Skin and Skin Equivalents With T Cells**

Normal healthy human skin obtained during plastic surgery was cut into small pieces (approximately 6 mm × 3 mm). This skin was then placed in the upper compartment of Transwell plates, not allowing cell-cell contact, and was exposed for 3 days to heterologous unstimulated and stimulated CD45RO+ T cells that were placed in the lower compartment (106/well). From normal, healthy human skin obtained during plastic surgery, a three-dimensional whole-skin model was built up from dermal fibroblasts and a biomatrix and multilayered epidermis with stratum corneum, according to the method of the reconstructed skin equivalent ast-2000 (CellSystems Biotechnologie Vertrieb GmbH, St. Katharinien, Germany) (27). The cultured skin equivalents, grown for 2 weeks at the air-liquid interface, were then exposed together with their inserts (not allowing cell-cell contact) for 3 days to heterologous unstimulated and stimulated CD45RO+ T cells. The skin pieces were then fixed in Bouin’s solution (Sigma Chemical Co.) and were further processed for staining with hematoxylin/eosin and TUNEL.

**Statistical Analysis**

Results are shown as mean ± SD. The paired Student’s t test was used for comparison of paired conditions.

**RESULTS**

**KC Injury Induced by Activated T Cells**

The histologic hallmark of eczematous disorders consists of a marked KC pathology. Acantholysis and spongiosis in the epidermis is characterized by impairment or loss of cohesion between KCs and an influx of fluid from the dermis, progressing to vesicle formation (Figure 1a1a and ref. 28). In preliminary experiments, we observed several morphologic changes in cultured primary human KCs (Figure 1b1b) 1b in the coculture with autologous T cells. Resting, unstimulated T cells did not affect KC morphology (Figure 1c1c), whereas KCs cocultured with stimulated T cells were shrunken and rounded; they finally detached from the plates and died (Figure 1d1d). The type of KC death was investigated by demonstration of apoptotic features such as chromatin condensation and fragmented nuclei made visible by HOECHST staining (26). To eliminate the possibility that the apoptotic bodies were derived from T cells, KCs and T cells were separated in Transwell plates. As shown in Figure 1f1f, KC apoptosis was induced by soluble factors from stimulated T cells, whereas resting T cells did not show any effect (Figure 1e1e). These results demonstrated that KCs are target cells of activated T cells that invade the epidermis in the effector phase of eczematous disorders, suggesting that KC injury and death may have a consistently contributory role to the KC pathology seen in eczema.
Role of IFN-γ and fas in KC Apoptosis

We then focused on the mechanism of KC death induced by activated T cells. FasR (CD95) is a 47-kDa membrane protein that transduces an apoptotic signal into the cell when triggered by its ligand (FasL) or by agonistic Abs (29, 30). KCs expressed detectable levels of FasR mRNA, which significantly increased after incubation with IFN-γ or with supernatants from stimulated CD45RO+ (memory/effecter) T cells (Figure 2a). In contrast, FasL mRNA was not detected in the KCs. As shown in Figure 2b, unstimulated KCs showing low FasR expression were resistant to Fas-mediated killing. IFN-γ, which is known to induce growth arrest and the expression of squamous differentiation genes (31), showed no significant killing activity. However, KCs pretreated with IFN-γ (but not TNF-α, IL-4, IL-5, IL-12, or IL-13; data not shown) were efficiently killed by cross-linking of FasR with anti-FasR mAb or soluble FasL (Figure 2a). In KCs, the sensitivity to Fas-induced death correlated very well with levels of cell-surface FasR (Figure 2c). Increasing doses of IFN-γ induced FasR expression on KCs. When the number of FasR molecules on KCs reached a threshold of approximately 40,000 per KC, the cells became susceptible to apoptosis.

Figure 2: (a) Expression of FasR mRNA by primary human KCs. Lane 1: unprotected template. Lane 2: after 8 hours of KC culture. Lane 3: after 24 hours of KC culture. Lane 4: after 16 hours of KC culture followed by 8 hours with 1.0 μg/mL anti-FasR mAb. Lane 5: after 16 hours of KC culture followed by 8 hours with diluted (50%) supernatant (sup.) from stimulated CD45RO+ T cells. Lane 6: after 16 hours of KC culture followed by 8 hours with 1.0 ng/mL IFN-γ. A representative result of three experiments is shown. (b) IFN-γ and Fas-induced KC apoptosis. KC viability was monitored by ethidium bromide exclusion and flow cytometry. Treatments shown are control (KCs alone), 1 μg/mL activating anti-FasR mAb, 10 ng/mL IFN-γ, and 1 μg/mL anti-FasR mAb added 1 day after starting incubation with 10 ng/mL IFN-γ. AP < 0.05. (c) IFN-γ-induced FasR counts exhibit a threshold for KC apoptosis. KCs were pretreated with the indicated doses of IFN-γ; 1 μg/mL anti-FasR mAb was added 1 day after starting incubation with IFN-γ. KC viability was assessed by ethidium bromide exclusion and flow cytometry. The threshold KC viability was monitored by ethidium bromide exclusion and flow cytometry. In the flow cytometry setting, KCs and T cells are gated according to forward and side scatter. Both cell populations were therefore monitored separately. Coculture of primary human KCs and autologous unstimulated or stimulated (with anti-CD2, anti-CD3, and anti-CD28 mAb) CD45RO+ T cells. AP < 0.05. Inhibition of CD45RO+ T-cell–induced KC death by 1 μg/mL blocking anti–IFN-γ receptor mAb and 20 μg/mL neutralizing anti–IFN-γ mAb. Results in b–d represent mean ± SD of triplicate cultures from three different experiments. Control, KCs alone.

Induction and Regulation of KC Apoptosis by T cells

Skin-infiltrating T cells belong primarily to the CD45RO+ subset (32, 33). In KC–T cell cocultures, we observed killing of KCs by stimulated autologous CD45RO+ T cells (Figure 2d; P < 0.05). Ligation of the T-cell receptor–CD3 complex by agonist peptides or mAbs to T-cell receptor–CD3 triggers a series of activation events, eventually leading to cell proliferation and cytokine production (34, 35). It is known that anti-CD3 mAb cross-linking induces the expression of cell-surface and soluble FasL by T cells (36). Accordingly, KC killing was significant only after activation of T cells, either with phorbol ester and calcium ionophore, cross-linking of the T-cell receptor–CD3 complex by plate-bound anti-CD3 (data not shown), or a combination of anti-CD2, anti-CD3, and anti-CD28 mAbs (Figure 2d; P < 0.05). Cytokine pretreatment of KCs was not necessary for death; however, it is apparent that sufficient IFN-γ was produced by the stimulated T cells themselves to upregulate KC FasR expression. As seen in Figure 2d, KC killing could be inhibited by pretreatment with either neutralizing anti–IFN-γ mAb or by the IFN-γ receptor–blocking mAb. Neutralizing mAb's against IL-4, IL-5, and IL-13, and the mutant IL-4–antagonist showed no inhibitory activity (data not shown). We further examined whether FasR engagement on IFN-γ–treated KCs triggered apoptotic cell death. Stimulated T cells induced KC apoptosis as demonstrated by HOECHST staining (Figure 2f,1), by annexin V staining of membrane phosphatidylserine inversion in KC–T cell cocultures (Figure 2a), and by DNA fragmentation in Transwell cocultures with no cell-cell contact allowed (Figure 2b). This demonstrates that soluble FasL released from T cells induces apoptosis in KCs in the absence of cell-cell contact. KC apoptosis was again inhibited by neutralizing IFN-γ, blocking the IFN-γ receptor, and by pretreatment of KCs with the caspase inhibitor Z-VAD-FMK (37). To further test that FasR expression on KCs is triggered by FasL from stimulated T cells, we used the human Fas-Fc protein as a competitive inhibitor of FasL–FasR interactions (38). We found that the Fas-Fc protein effectively inhibited apoptosis of KCs induced by stimulated T cells (Figure 3a, a and b).

Figure 3: Induction and regulation of KC apoptosis in KC–T cell cocultures, (a) Coculture of primary human KCs and stimulated autologous CD45RO+ T cells. After 3 days of coculture, cells were stained with annexin V and subjected to flow cytometry. With the indicated stimuli, the number of apoptotic KCs increases and the cells show an increase in annexin V binding (open histograms). Filled histograms show live KCs. The percentage of apoptotic KCs is indicated at upper right. Panel 1: KC apoptosis induced by coculture with CD45RO+ T cells stimulated with anti-CD2, anti-CD3, and anti-CD28 mAb's. Panel 2: Isotype control mAb. Panel 3: Blocking anti–IFN-γ receptor mAb (1 μg/mL). Panel 4: Fas-Fc protein (1 μg/mL). Panel 5: The caspase inhibitor Z-VAD-FMK (50 μM). Panel 6: Z-VAD-FMK (100 μM). (b) Coculture of primary human KCs and stimulated autologous CD45RO+ T cells in Transwell plates. After 3 days in coculture, permeabilized KCs were stained with propidium iodide and subjected to flow cytometry. Filled histograms demonstrate control KCs containing diploid DNA. The percentage of apoptotic KCs with hypodiploid DNA (open histograms) is shown in the upper right corner. Panel 1: Primary human KCs alone were cultured in the lower well. Panel 2: KCs were pretreated with 10 ng/mL IFN-γ for 24 hours, and KC apoptosis was determined 3 days after 1 μg/mL activating anti-FasR mAb was added. Panel 3: KC apoptosis induced by Transwell coculture with stimulated CD45RO+ T cells. Panel 4: Inhibition of KC apoptosis induced by IFN-γ and anti-FasR mAb, with 1 μg/mL ZVAD blocking mAb. Panels...
5 and 6: Inhibition of KC apoptosis induced by Transwell coculture with stimulated CD45RO+ T cells, with 1 μg/mL Fas-Fc protein (panel 5), and with 10 μg/mL neutralizing anti–IFN-γ mAb (panel 6). Results in a and b are representative of three experiments.

**Induction of KC Apoptosis by Type 1 and Type 2 T Cells**

To establish a more disease-related model for KC apoptosis in eczematous disorders, we investigated the effects of different phenotypes of stimulated T cells on KCs. Type 1 CD4+ and CD8+ T cells contain and secrete primarily the cytokine IFN-γ, and only low amounts of IL-4 and IL-5. In contrast, type 2 CD4+ and CD8+ T cells contain and secrete IL-4, IL-5, and IL-13, but only low amounts of IFN-γ (39). The type 1 CD4+ and CD8+ T cells were able to induce KC death (Figure 4a; P < 0.05). In contrast, type 2 T cells had no effect on KCs after differentiation. Additionally, we combined type 1 and type 2 T cells and measured KC apoptosis. This mixture induced apoptosis less than type 1 cells alone did, but apoptosis was still highly significant (data not shown). IL-12 is produced mainly by macrophages and dendritic cells, and promotes type 1 T-cell responses by selective upregulation of IFN-γ production by T cells (40). Stimulation of type 2 CD4+ or CD8+ T cells with IL-12 increased IFN-γ production and ablated IL-4 production (Figure 4b; P < 0.05). In this case, IL-12 stimulation of type 2 T cells enabled the induction of KC apoptosis (Figure 4c; P < 0.05). Supernatants from stimulated type 1 and type 2 T cells released soluble FasL; addition of IL-12 did not influence the secretion of soluble FasL (Figure 4d).

**Features of Inflammation in Eczematous Dermatitis**

The histologic appearance of acute lesional AD and ACD skin was defined by various degrees of dermal perivascular infiltration with mononuclear cells, mostly consisting of CD4+ T cells as well as CD8+ T cells (Figure 5a–c). The expression of CD45RO on the T cells suggests a previous encounter with antigen; many of the T cells showed signs of intraläsional activation as defined by membrane expression of HLA-DR (2). It should be noted here that T cells constitute the major cells in dermal infiltrates, and that some of the CD4+ and CD8+ T cells invade the epidermis (Figure 5b and c). Therefore, membrane-bound FasL on T cells can trigger FasR on KCs by cell-cell contact. Figure 5d demonstrates FasL immunoreactivity on lesional T cells of acute AD. The basal and suprabasal compartment of the epidermis was strongly reactive for FasR, and the KCs showed high FasR surface expression in acute AD (Figure 5e). Similarly, ACD skin showed FasL immunoreactivity on KCs (data not shown). Chronic AD lesions displayed weaker FasR immunoreactivity in basal KCs. Healthy control skin displayed no inflammatory infiltrate and no KC staining with anti-FasR mAb. KCs from healthy individuals and from AD and ACD patients expressed IFN-γ receptors on their surface in vitro and in vivo (Figure 5f and ref. 41). In Figure 6, c and d, condensed and fragmented nuclei of apoptotic cells are demonstrated in acute AD, chronic AD, atopy patch tests, and acute ACD, we also investigated apoptosis in cocultures of normal human skin of healthy individuals with stimulated T cells, and in cocultures of skin equivalents with stimulated T cells.
Figure 5: Features of inflammation in AD. Immunohistologic staining with 3-amino-9-ethylcarbazole substrate and counterstaining with hematoxylin. (a) Subepidermal inflammatory infiltrate, consisting mainly of lymphocytes. ×100. (b) CD4+ cells infiltrating the spongiotic epidermis. ×200. (c) CD8+ cells. ×200. (d) FasL+ cells in the dermal cellular infiltrate. ×200. (e) Strong immunoreactivity of FasR in the basal epidermis; weaker staining in the T-cell infiltrate. ×100. (f) Detection of IFN-γ receptor on KCs and infiltrating T cells. ×200.

Demonstration of KC apoptosis in AD, atopy patch tests, ACD, and cultures of healthy human skin and skin equivalents.

Skin biopsies from acute AD, chronic AD, atopy patch tests, and acute ACD were analyzed using the TUNEL technique (Figure 6, a and b) and HOECHST staining (Figure 6, c and d). Patch-test reactions to aeroallergens constitute a suitable model for the study of allergic inflammation in AD. They are specific for sensitized AD patients, and do not occur in healthy volunteers or in patients suffering from asthma or rhinitis (42). As a negative control, we used normal skin from healthy individuals. No TUNEL-stained cells were observed in the stratum basale or stratum spinosum of normal skin. TUNEL-stained cells were observed in the stratum basale or stratum spinosum of normal skin. TUNEL-stained nuclei were rarely seen in the lumen of spongiotic vesicles of acute AD and ACD. As shown in Figure 6, a and b, TUNEL-stained KCs are visible primarily in the cohesive epidermis (stratum spinosum) of acute lesions in which acantholysis and spongiosis had not progressed to vesicle formation. In chronic AD, TUNEL-stained KCs are visible at a lower frequency than in acute AD and ACD. HOECHST staining was used as a second method, complementing TUNEL staining.

DISCUSSION

This study demonstrates that Fas-induced KC apoptosis caused by skin-infiltrating T cells is a major mechanism in the pathogenesis of eczematous dermatitis. In recent years, much interest has focused on apoptosis as a mechanism to control cell numbers (44). The interactions between FasL and FasR are important in maintaining lymphoid cell numbers. However, it is likely that FasL-FasR interaction also plays a role in other important biologic processes. Various cells in the body constitutively express FasR and can be induced to do so (45, 46), and the sensitivity to Fas-induced KC death correlated very well with numbers of cell surface FasR in our study. An explanation for this may be that quantitative increases in FasR expression above a certain threshold level result in a signaling intensity that promotes apoptosis. On the other hand, it is possible that the signal transduction pathway for Fas-induced apoptosis is functionally inactive in unstimulated KCs. The threshold for IFN-γ in the induc
tion of FasR on KCs appears to be in the range of 0.1–1.0 ng/mL.

FasL expression is usually limited to activated T cells, natural killer cells, and cells of certain immunologically privileged sites (49). T cells appear to be important players regulating inflammatory processes in eczematous dermatitis (6–18). This is supported by the observation that immunosuppressive drugs such as FK-506 and glucocorticoids, which block T-cell activation, are effective in treatment of eczematous disorders (4, 5). The coupling of Fas signaling to the death pathway involves receptor trimerization followed by the binding of adapter proteins and then the caspase enzyme cascade, ultimately resulting in apoptosis (29, 30). Our studies using the caspase inhibitor Z-VAD-FMK (37) show that caspase activation is also critical for Fas-mediated apoptosis of KCs. In addition, recombinant Fas-Fc protein (38) prevents T cell-mediated KC apoptosis. This protective effect indicates that the Fas-L:FasR interaction is responsible for T cell–mediated KC apoptosis. The lethal hit is delivered to KCs by Fasl, expressed on the surface of T cells that invade the epidermis, and probably by soluble Fasl, released from T cells. IFN-γ, which increases FasR expression on KCs, is a further prerequisite in T cell–mediated KC apoptosis. This is demonstrated by the fact that blocking of IFN-γ totally abrogates KC apoptosis induced by activated T cells. Partial contributions by other cytokines and apoptotic mechanisms were principally eliminated in experiments demonstrating neutralization of the effects by blocking IFN-γ and by using Fas-pathway antagonists. It has been proposed that the basal epidermis is resistant to apoptosis due to highly developed antiapoptotic defenses, which are reversible and decrease during differentiation (26). Adding IFN-γ causes cultured KCs to differentiate and lose resistance to apoptosis (31). Thus the increased susceptibility to killing by Fas ligation in vitro may be due to both increased FasR expression and reduced antiapoptotic defenses.

Eczematous skin lesions with distinct etiology are associated with T-cell infiltration in the epidermis, contact between T cells and KCs, and a marked KC pathology (28, 50, 51). KCs in inflamed skin express HLA-DR and ICAM-1, which are not expressed on KCs of normal skin (1, 2). The only mediator known to date that induces HLA-DR on KCs is IFN-γ. Together, these data suggest a model of T cell–induced KC apoptosis in the pathogenesis of eczematous dermatitis. In an early step, T cells infiltrating the epidermis attach to KCs. Subsequently, the apoptosis signal is delivered to KCs by Fasl expressed on the surface of T cells. In our experiments, Fasl was detected only on the T-cell surface — not on KCs in AD, ACD, and patch-test lesions. However, in a study of toxic epidermal necrolysis, FasL expression on highly activated KCs was shown (52). In addition to Fasl:FasR interaction, it is possible that activated T cells in eczematous dermatitis use granule-mediated killing by perforin and granzymes (53–55). However, granule-mediated cytotoxicity is not blocked by soluble Fas-Fc protein. Additionally, using immunohistology, we could not detect perforin or granzyme B in lesional skin of ACD and AD patients.

To establish a more disease-related model for KC apoptosis in eczematous dermatitis, we investigated the effects of different subsets of activated T cells on KCs. We demonstrated Fas-mediated apoptosis by both CD4+ and CD8+ type 1 T cells. There is accumulating evidence that not only CD4+, but also CD8+ T cells possess the potential to both stimulate IgE production and promote allergic inflammation (10, 56). AD is regarded as a type 1 T cell–mediated phenomenon in vivo, based on a number of observations (6, 7). Several reports suggest that type 1 and type 2 cytokines secreted from activated T cells are important for the pathogenesis of AD (17). Clinical and histologic features of AD are almost indistinguishable from those present in ACD. Recent studies found IFN-γ mRNA and protein highly expressed in eczematous skin of the vast majority of AD patients (15). IL-12 stimulation of type 2 T cells abolished IL-4 and induced IFN-γ production without major changes in IL-5 and IL-13 production. This explains the biphasic nature of the cytokine response observed in acute versus chronic AD lesions. Acute lesions and early response after epidermal allergen application were characterized with a predominant type 2 cytokine pattern (4, 16, 57). Increased levels of IL-12 mRNA was demonstrated in chronic AD lesions (58), and chronic lesions were characterized by increased IFN-γ and decreased IL-4 levels, with more or less constant IL-5 and IL-13 levels (13, 15, 16, 57, 58). We observed killing of KCs by type 2 T cells after stimulation in the presence of IL-12. A cause-and-effect relationship between IL-12 and IFN-γ expression may play a role in the conversion of type 2 T cells into IFN-γ–producing T cells that induce KC apoptosis in AD (15–17, 58). In addition, virus infections and bacterial superantigens of AD skin may lead to IFN-γ secretion of skin-infiltrating T cells (14, 58, 59). Furthermore, KCs exhibited a relatively low threshold for FasR expression in response to IFN-γ, in the range of 0.1–1.0 ng/mL.

Our results suggest that KC apoptosis is the initiating event in the development of the epidermal pathology seen in eczematous dermatitis. TUNEL and HOECHST staining showed that apoptotic KCs were present in the epidermis of patients with eczematous disorders. Most notably, KC apoptosis occurs in suprabasal cells, where spongiosis and acantholysis takes place. The basal layers of lesional epidermis had high levels of FasR, but it is likely that antiapoptotic mechanisms prevent apoptosis of basal KCs (26). Apoptosis of individual KCs is the first event leading to disruption of epidermal continuity and vesicle formation. Damage to KCs leads to the loss of intercellular cohesion (acantholysis) and subsequent cell detachment. Fluid influx from the dermis and intercellular edema contributes to spongiosis, which is defined as widening of the intercellular space and a spongeliike appearance of the epidermis (28). Based on our findings, the reason that not all the suprabasal KCs that express FasR undergo apoptosis uniformly in situ has to do with the activation of KCs induced by IFN-γ and the ligation of FasR on the KCs by Fasl expressed on T cells infiltrating the epidermis. Apoptosis of epidermal KCs is known to be an important feature in lichenoid tissue reaction, fixed drug eruption, graft-versus-host reaction, and the effects of ultraviolet radiation (26, 60, 61).

Skin equivalents formed by KCs obtained from healthy skin and individuals with no skin disorders that are cocultured with fibroblasts embedded in collagen lattices represent promising tools for studies of skin pathology (27, 62). In cultured skin equivalents, KC differentiation and composition of the basement membrane are not exactly the same as in living epidermis. Therefore it is likely that the pro- and antiapoptotic mechanisms operating in KCs are somewhat different in vivo, and basal KCs are probably more sensitive to apoptosis in skin equivalents. We also used the short-term culture of normal human skin in a medium maintaining KC viability to investigate the effects of activated T cells on epidermal KCs. The induction of KC apoptosis in normal human skin and cultured skin equivalents after exposure to activated T cells establishes the in vivo relevance of T cell–mediated, Fas-induced KC apoptosis.

The knowledge of this molecular basis is pivotal in understanding the development of pathology in eczematous disorders, and opens a future for more focused therapeutic applications.

ACKNOWLEDGMENTS

This work was supported by grants from the Deutsche Forschungsgemeinschaft (TR460/1–1), the Swiss National Foundation (31.50590.97/1), and the Baumgarten Foundation (Zürich). We thank A. Speiser (Landspital Davos), J. Kammerer (Department of Pathology, Rätisches Kantonsspital Chur), R. Gillitzer, W. Dummer, A. Toksoy, H. Krenig, and C. Siedel (Department of Dermatology, University of Würzburg) for their work.

REFERENCES

3. Dustin ML, Singer KH, Tuck DT, Springer TH. Adhesion of