Effect of the Monocyte Locomotion Inhibitory Factor (MLIF) a Natural Anti-Inflammatory Produced by E. histolytica on Apoptosis in Human CD4+ T Lymphocytes

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ABSTRACT

Apoptosis prevents the extravasation of intracellular material and the subsequent inflammatory response. Currently, it is not known whether Monocyte Locomotion Inhibitor Factor (MLIF), an anti-inflammatory pentapeptide, induces programmed cell death. We evaluated the effect of MLIF on extrinsic and intrinsic apoptosis pathways human CD4+ T lymphocytes. Cells were cultured for 24 h in RPMI-1640 medium alone (control) or in RPMI medium containing MLIF alone, PMA alone, PMA + MLIF or actinomycin D. Annexin V/propidium iodide-stained cells in early apoptosis showed that cells treated with MLIF or PMA + MLIF were not significantly different from control cells in medium; in contrast, cells treated with PMA or PMA + MLIF demonstrated significant differences from the control in delayed apoptosis. Cytochrome c and caspase 3 levels in cells treated with MLIF showed no significant differences from control cells, however, compared to the control, cells treated with PMA and PMA + MLIF exhibited a significant increase in cytochrome c and caspase 3 levels, which demonstrates that this weak induction of cell death is regulated by the intrinsic pathway of apoptosis. The Fas receptor was not detected in cell culture with any of the treatments employed, suggesting that the extrinsic pathway of apoptosis is not involved. The MLIF per se does not induce apoptosis in human CD4+ T lymphocytes; there may be an additional effect of PMA + MLIF producing the low levels of cell death recorded in the late apoptosis phase. MLIF acts as a natural, biological anti-inflammatory compound produced in axenic cultures of E. histolytica that does not cause apoptosis or elicit an immune response due to its small size, which could make it a strong candidate for future clinical applications.

Keywords: MLIF, Apoptosis, Anti-Inflammatory, Intrinsic Pathway, Extrinsic Pathway

1. Introduction

T lymphocytes constitute an essential part of the immune system. Their generation, activation, proliferation and survival are subject to tight regulation by several extracellular factors including cytokines, MHC-antigen complexes and co-stimulatory ligands; however, internal or external stimuli, such as over-stimulation, cytotoxicity or time, can lead to death. The balanced interplay between these factors determines the fate of a T cell [1-3]. Once the cytotoxicity or over-stimulation occurs, T cells become apoptosis-sensitive, leading to the death of most cells and the survival of only a few memory T cells. Like other cells, T cells are dependent on signals from their environment for their survival; apoptosis thus represents a tightly regulated process by which lymphocyte homeostasis is controlled, and the generation of autoreactive cells is prevented [4-6].

Apoptosis or programmed cell death, which plays a central role in the normal development of multicellular organisms, is essential for T cell function, the regulation of B cells, the suppression of autoimmunity, the control infections, as well as immune surveillance and homeostasis [7,8]. One of the most important physiological characteristics of cell death by apoptosis is no inflammatory response. Key proteins are released into the cytosol by mitochondria (intrinsic pathway of apoptosis) or as a
consequence of activating a cell death receptor on the cell surface (extrinsic pathway of apoptosis) [9]. Both signalling pathways can converge upon the activation cascade of effector caspases. Activation of the pro-apoptotic proteins of the Bcl-2 family leads to the formation of a pore in the outer membrane of mitochondria, which allows the release of proteins of including cytochrome c from mitochondrial intramembranous space into the cytosol; activates the apoptosome complex and caspase 9; and finally activates effector caspases, such as caspase-3-6 and 7 which triggers the final stages of apoptosis. In contrast, when a cell dies via necrosis, the cytoplasmic contents are released into the interstitial space, exposing a large number of antigens and triggering a strong inflammatory response [10,11].

The protozoan parasite _Entamoeba histolytica_ (E. histolytica) produces an anti-inflammatory pentapeptide (Met-Gln-Cys-Asn-Ser; MLIF) that inhibits human monocyte locomotion in _vivo_ and _in vitro_ [12]. MLIF depresses production of nitric oxide reactive intermediates by human monocytes and neutrophils, decreases the expression of adhesion molecules (VL-4 in monocytes and VCAM-1 in the vascular endothelium), and inhibits delayed hypersensitivity reactions in gerbils and guinea pigs [13-15]. The scrambled pentapeptide (MLIF, Gln-Cys-Met-Ser-Asn); showed no anti-inflammatory properties [16]. This anti-inflammatory effect could be attributed to the chemical activity of the peptide. Ongoing studies in quantum chemistry have revealed that a pharmacophore group in the MLIF sequence (… Cys-Asn-Ser) could be responsible for most of the anti-inflammatory properties of this molecule [17]. Currently it is unknown whether MLIF may have some involvement with the mechanisms of programmed cell death. The aim of this study was to evaluate the effect of anti-inflammatory MLIF on apoptosis _in vitro_ by studying well-characterized markers of apoptosis, such as annexin-V/propidium iodide (IP) binding, activation of caspase-3, cytochrome c and Fas receptor expression in human CD4+ T lymphocytes.

### 2. Materials and Methods

#### 2.1. Monocyte Locomotion Inhibitory Factor (MLIF)

MLIF (Met-Gln-Cys-Asn-Ser), 96% pure was obtained commercially (American Peptide Co., Sunnyvale, CA, USA). All working solutions were tested for endotoxin (LPS < 0.3 pg) using the limulus assay (Amoebocyte Lysate Endosafe KTA Charles River Endosafe INC, Charleston, Inc, USA) and were kept at −70°C until use.

#### 2.2. Cell Purification

Thirty milliliters of heparinized venous blood was obtained from healthy, non-smoking adult volunteers of both sexes. Blood was diluted 1:2 with phosphate buffered saline (PBS; 0.15 M phosphate buffer), then 10 ml of the sample was layered over 4 ml of a Ficoll-Hypaque (Sigma Chemical Co., St. Louis, MO) gradient (δ = 1.077) and centrifuged at 700 × g for 30 min at 21°C [18]. Peripheral blood mononuclear cells (PBMC) at the interface were removed and washed twice with PBS. CD4+ T cells were purified using the CD4+ T Cell Isolation Kit II (Miltenyi Biotec, Germany) an indirect magnetic labeling system for the isolation of untouched CD4+ T cells from human PBMCs, with LS columns and a MidiMACS separator. Briefly, 1 × 10⁷ PBMC were placed in polypropylene tubes with 80 μl of PBS-albumin-EDTA and 20 μl a cocktail of biotin-conjugated antibodies against CD8, CD11b, CD16, CD19, CD36, CD56, CD123, TCRγδ and CD235a (glycophorin A), and were incubated for 10 min at 4°C. These cells were subsequently magnetically labelled with Anti-Biotin MicroBeads for depletion. The population of CD4+ T lymphocytes that was isolated was 95% pure.

#### 2.3. Cell Culture

Five × 10⁵ CD4+ T cells were placed in 24-well plates in:

1) RPMI-1640 medium alone (supplemented with 10% fetal calf serum (FCS), 2 mM L-glutamine, 100 U/ml streptomycin, 5 μg/ml gentamicin, and 1 mM sodium pyruvate) (Gibco Laboratories, Grand Island, NY, USA);
2) in medium supplemented with 50 ng/ml phorbol 12-myristate 13-acetate (PMA) (Sigma, Chemical Co, St. Louis MO); 3) in medium with 50 μg/ml phorbol myristate acetate (PMA) (Sigma, Chemical Co, St. Louis MO); 4) PMA + MLIF; or 5) 1μg/ml actinomycin D for 24 h at 37°C, 5% CO₂ [19]. Cell viability was ≥90% by trypan blue dye (Sigma) exclusion. The optimal concentrations of MLIF and PMA were determined from dose-response curves.

#### 2.4. Annexin V/Propidium Iodide Double Staining by Flow Cytometry

For annexin V/propidium iodide (PI) double staining, cells were resuspended in 100 ml of binding buffer (10 mM N-(2-hydroxyethyl) piperazine-N-2-ethanesulfonic acid (HEPES)/NaOH, 140 mM NaCl, 2.5 mM CaCl₂, pH 7.4) and stained with 5 ml of both FITC-conjugated annexin V and PI (BD Biosciences). The mixture was incubated for 15 min at room temperature in the dark, followed by flow cytometry analysis. Acquisition of 10,000 events was conducted in flow cytometry FACScalibur (BD Biosciences, Palo Alto, CA). For analysis, Win MDI
2.8 software was used.

2.5. Quantitation of Total Cytochrome c in Cultured Cells

After a 24 h incubation in the presence of medium alone, PMA, PMA + MLIF or actinomycin D the CD4+ T cells were harvested in their culture medium, pelleted by centrifugation at 300 × g for 15 min, rinsed with PBS, and resuspended in Cell Lysis Buffer 2 (Cell Lysis Buffer 2 solubilises the cytochrome c in both the cytosol and mitochondria) to a final concentration of 1.5 × 10^6 cells/ml. The cell lysate was incubated for 1 h and centrifuged at 1000 × g for 15 min. The resulting supernatant cytochrome c was measured with a quantitative sandwich enzyme immunoassay technique (cytochrome c ELISA Kit) (Calbiochem® Merck KGaA, Darmstadt, Germany). Plates were pre-coated with a monoclonal antibody specific for cytochrome c. Standards and samples were pipetted into coated wells, and cytochrome c was bound to the immobilised antibody. After washing, an enzyme-linked monoclonal antibody specific for cytochrome c was added to each well, followed by the substrate. In this assay, the colour develops in proportion to the amount of cytochrome c bound in the initial step. Colour development was stopped, and the intensity of the colour was measured. The quantity of human cytochrome c was determined as a change in absorbance at 450 nm using a microplate reader (Dynatech MR 5000, Maryland, USA).

2.6. Quantitation of Activated Human Caspase-3

Caspase-3 activity was measured by using a human active caspase 3 ELISA kit protocol (OptEIA™, BD Biosciences). Briefly, cells were lysed in a hypotonic buffer (10 mM HEPES [pH 7.4], 50 mM NaCl, 2 mM MgCl₂, 5 mM EGTA, 1 mM phenylmethylsulfonyl fluoride, and protease inhibitor cocktail). After incubating for 45 min on ice, lysates were centrifuged at 40,000 × g for 45 min at 4°C. Cell lysates were diluted with 200 μl assay diluent buffer (50 mM HEPES [pH 7.4], 0.2% CHAPS {3-[(3-cholamidopropyl)-dimethylammonio]-1-propanesulfonate}, 20% glycerol, 2 mM EDTA, 10 mM dithiothreitol), was added to each well and incubated for 2 h at room temperature. Next, 100 μl of cytochrome c standard or sample was added to each well and incubated for 2 h. Subsequently, 100 μl of horseradish peroxidase-conjugated streptavidin (1X) and 100 μl of substrate solution were added to each well and incubated for 30 min. Finally, 50 μl of stop solution was added to each well. The absorbance of the cleaved substrate at 450 nm was measured with a microplate reader (Dynatech MR 5000, Maryland, USA).

2.7. Quantification of Fas in the Supernatant of Cultured Cells

The presence of Fas in the supernatant fluids of cultured CD4+ T lymphocytes (1 × 10^6 cells/well) was determined and quantified using an ELISA kit with an antibody that recognises human Fas (Ray Biotech, Inc., Atlanta, Georgia), according to the manufacturer’s instructions. For calibration, we used a commercially available, natural human Fas protein to construct a standard curve and to obtain absolute values. The concentration of Fas was measured in six independent assays per treatment, each in duplicate, and the average of the six measurements was considered to be the final concentration.

2.8. Statistical Analysis

All values were expressed as mean ± standard deviation (SD). Commercial SPSS v. 11.0 for Windows (SPSS Inc., Chicago, IL) was used for statistical analysis. Statistical comparisons between groups were performed using the Mann-Whitney U tests; p values ≤ 0.05 were deemed statistically significant.

3. Results

3.1. Detection of Apoptosis by Annexin V/Propidium Iodide Staining

To determine whether apoptosis was induced by MLIF, and to discriminate between early apoptosis and necrosis, cells were simultaneously stained with annexin V and PI. Human CD4+ T lymphocytes were cultured in the presence of RPMI medium (negative control), MLIF, MLIF, PMA or PMA + MLIF, and actinomycin D served as the positive control. Results showed that 7% of CD4+ T lymphocytes underwent spontaneous apoptosis in the medium control. When the cells were stimulated with MLIF, 9% of CD4+ T lymphocytes stained positive for annexin V. When stimulated with PMA, a significant number of cells (13%) were apoptotic (p < 0.003 compared to medium control), and among cells treated with PMA+MLIF, the frequency of apoptosis was 10% (not significant, NS). Actinomycin D treatment increased early apoptosis to 19% of cells (p < 0.0001 compared to medium control) (Figure 1). However, in the analysis of late apoptosis, PMA + MLIF significantly increased the binding of annexin V (21% of cells) compared to the control (11%, p < 0.002) (Figure 1), suggesting that the pro-apoptotic activity of MLIF + PMA is related to the extrinsic signalling pathway and/or the intrinsic pathway of apoptosis.

3.2. Production of Cytochrome c and Fas (APO-1)

PMA + MLIF may favour activation-induced cell death.
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Figure 1. Apoptosis in CD4+ T lymphocytes. Lymphocytes were cultured for 24 h in RPMI-1640 medium alone (control) or in RPMI medium containing MLIF alone, PMA alone, PMA + MLIF or actinomycin D. Cells were stained with FITC-conjugated annexin V and propidium iodide (annexin/IP) and processed for cell death determination by flow cytometry. (a) Calibration parameters and positive control (actinomycin D); (b) The numbers in each quadrant of the flow cytometry histograms represent the mean percentage of six independent experiments. The percentage of necrotic cells is indicated in the upper left quadrant; apoptotic-necrotic cells, in the upper right quadrant; and early apoptotic cells in the lower right quadrant.

Non-stimulated CD4+ T cells (RPMI medium control) produced 9 μg/ml of cytochrome c; cells stimulated with MLIF produced 12 μg/ml; with PMA, 15 μg/ml; with PMA + MLIF, 19 μg/ml; and with actinomycin D, 33 μg/ml (Figure 2). MLIF per se did not stimulate the release cytochrome c (NS), but treatment with PMA + MLIF resulted in a there was a statistically significant difference compared to the negative control (p < 0.001) and positive control (p < 0.0001), suggesting that the intrinsic pathway of apoptosis is activated. Actinomycin D produced a greater increase in the release of cytochrome c (Figure 2). In contrast, Fas receptor expression was not observed under any of the treatment conditions (data not shown).

3.3. Determination of Activated Human Caspase-3

Execution of apoptosis requires the activation of the caspase cascade, where activated initiator caspases cleave and thereby trigger downstream effector caspases that, in turn, dismantle the cell. Whereas proteolytic cleavage of an effector caspase is indicative of its activation, the same is not necessarily true with initiator caspases.

We wondered whether the effect of MLIF on the expression of cytochrome c converged upon the caspase-3 pathway. In cells treated with medium alone, basal expression of caspase-3 was 90 U/ml (Figure 3). Exposure to MLIF increased expression of caspase-3 to 110 U/ml, and upon PMA treatment, was 196 U/ml. In cells treated with PMA + MLIF caspase-3 production was 235 U/ml, and with actinomycin D, it was 329 U/ml. As expected, actinomycin D treatment was significantly different (p < 0.002) compared to basal levels (Figure 3) and cells treated with MLIF, PMA, PMA + MLIF (p < 0.001, p < 0.0001 and p < 0.01 respectively). The signalling cascade of caspase 3 was different between groups, and this stronger between the PMA and actinomycin D.

4. Discussion

Apoptosis can be induced either as a consequence of key
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Figure 2. Quantification of cytochrome c. One × 10⁵ CD4+ T lymphocytes, cultured in the presence of PMA, MLIF or PMA + MLIF for 24 h, were lysed and centrifuged at 4000 × g, and cytochrome c in the supernatant was measured. Histograms show mean values ± SD of six independent experiments. Statistically significant differences between all groups and actinomycin D are indicated with an asterisk, p < 0.05 (Mann-Whitney test).

Figure 3. Quantification of caspase-3. After culturing under the indicated conditions, 10⁵ lymphocytes were lysed and centrifuged at 4000 × g. Released caspase-3 in the supernatant was measured using a commercially available enzyme-linked immunosorbent assay (ELISA) kit. The levels of caspase-3 are represented as means of caspase-3 concentrations vs the negative control group, or the positive control group (actinomycin D) in cells treated with MLIF, PMA, PMA + MLIF (p < 0.001, p < 0.0001 and p < 0.01 respectively). NS = no significant difference, n = 6, p < 0.05 (Mann-Whitney Test).

proteins being released into the cytosol by the mitochondria or by the activation of a death receptor on the cell surface. Both signalling pathways converge at the level of effector caspases, without causing the inflammation produced by necrotic cell death [20].

Infection with *E. histolytica* is associated with an acute inflammatory response [21]. However, it is not completely clear how *E. histolytica* modulates the pro-inflammatory response of the host. Previous work studying the effect of MLIF on inflammation, specifically the inhibition of interleukins, such as IL-2, IFN-γ and IL-1β, indicates that MLIF acts selectively in different cell lineages. At least two possible explanations could account for this effect: MLIF may either inhibit soluble pro-inflammatory or regulate apoptosis mediators [22, 23].

In this study, we determined that MLIF *per se* does not induce programmed cell death in human CD4+ T lymphocytes, as demonstrated in early apoptosis. Cell death was slightly increased in the presence of PMA (MLIF + PMA) but was detected only in late apoptosis (apoptosis/necrosis). MLIF has powerful but selective anti-inflammatory properties; this peptide inhibits locomotion (random, chemotactic and chemokinetic) of normal human peripheral blood mononuclear phagocytes in response to various chemoattractants, such as C5a-desArg, lym-
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phokine chemotactic or lymphocyte-derived chemotactic factor (LDCF), but has no effect on polymorphonuclear neutrophils [12]. MLIF is capable of downregulating some pro-inflammatory cytokines, including IFN-γ, IL-2, and IL-6, and of inducing the anti-inflammatory cytokines IL-4 and IL-10 [22]. MLIF delays the arrival of mononuclear leukocytes into Reuck chambers applied to the skin of healthy human volunteers [12]. In this study, the combination of PMA + MLIF caused an increased susceptibility to late apoptosis in lymphocytes, implicating the intrinsic apoptosis signalling pathway. This increase may illustrate an additive effect because PMA-dependent apoptosis was also slightly increased in the late apoptosis. The intracellular apoptosis machinery depends on caspases, which are proteases responsible for executing programmed cell death. Once activated, a caspase will in turn activate the next pro-caspase downstream, creating a proteolytic amplification cascade. Some of the activated caspases cleave other key proteins in the cell, such as those that form the nuclear lamina and the cytoskeleton, leading to the destruction of the nuclear framework, and activate DNases that degrade nuclear DNA. These changes underlie the nuclear and cytoplasmic structural modifications that occur in apoptotic cells [24,25]. Release of cytochrome c from mitochondria is an important step in mitochondria-mediated apoptosis; it leads to the activation of caspase 9 and then caspase 3 [26]. The fact that the Fas receptor was absent from the surface of MLIF-treated T cells suggests that there was no death receptor activation. Although Fas is mostly thought of as a death receptor that is capable of inducing the apoptosis of activated T cells via the caspase cascade [27], this receptor has also been implicated in effective T cell activation. Fas could act as a receptor with functional pleiotropy in models of T cell [28]. MLIF interacts with human leukocytes through a mannose-containing receptor [29], and it causes an increase in the number of pericentriolar microtubules and in the cAMP concentration, without a concomitant reduction in cGMP [30]. In U-937 cells, MLIF inhibits the expression of inflammatory cytokines, such as MIP-1α and MIP-1β, which are NF-κB pathway-regulated proteins, as was described by Utrera-Barillas [31]. The p65-p50 heterodimer comprises the most abundant form of NF-κB in a PMA-induced system. Temporary studies showed that MLIF induces p50 translocation, which could be due to the fact that MLIF stimulates cAMP synthesis and phosphorylation of NF-κB by protein kinase A, phosphorylation of IκB and subsequent NF-κB translocation [32]. MLIF strongly inhibits inflammation but does not induce early apoptosis, downregulates the Fas receptor, activates caspase 3 or trigger the release of cytochrome c, indicating that it is largely non-cytotoxic, however; we must be cautious in extrapolating these results to other population of lymphocytes such as CD8+ T cells and regulatory T cells.

We propose that this is a novel natural anti-inflammatory molecule produced in axenic cultures of E. histolytica that, owing to its small size (five peptides), is non-antigenic, thus making it a strong candidate for future clinical applications.

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