Rho GTPase Is Activated by Cytotoxic Necrotizing Factor 1 in Peripheral Blood T Lymphocytes: Potential Cytotoxicity for Intestinal Epithelial Cells

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Bacterial colitis is characterized by an acute inflammation with polymorphonuclear leukocyte (PMNL) migration into the digestive lumen in response to inflammatory cytokines (16, 22, 50). This transmepithelial migration of PMNL and intestinal epithelial cells have been extensively studied using an in vitro model (28, 33, 36). T lymphocytes play an essential role in the immune surveillance at the intestinal epithelium level (46). Dysregulated T-cell responses to enteric bacteria have been implicated in the pathogenesis of some models of colitis (29, 30). Nevertheless, the role of T lymphocytes during the acute phase of bacterial colitis has been poorly investigated.

The recruitment of T lymphocytes from the circulation into the mucosa requires the extravasation of these leukocytes from the microvasculature and their subsequent homing into the lamina propria (6). Leukocyte migration in response to chemoattractants is a crucial phenomenon in both the immune and the inflammatory response. Several members of the Rho family of small GTPases, named Rho, Rac, and Cdc42, act as key regulators of the actin cytoskeleton and different data suggest that Rho proteins are involved in the leukocyte adhesion through integrins activated by chemoattractants (32). Moreover, small GTPases can regulate T-cell chemotaxis in response to stromal cell-derived factor 1α (SDF-1α) (13). Indeed, overexpression of dominant negative Cdc42 and activated mutants of all these Rho GTPases inhibited SDF-1α-induced T-cell chemotaxis (13). Moreover, D’Souza-Schorey et al. have shown that Rac may increase integrin-mediated adhesion of T lymphocytes (14).

Certain strains of Escherichia coli produce a toxin named cytotoxic necrotizing factor 1 (CNF-1) (3, 4, 8). These strains can induce urinary or gastrointestinal tract infections (4, 15). CNF-1 mediates its effects via the permanent activation of small Rho GTP-binding proteins, by causing deamidation of p21 Rho Glu63 residue (5, 19–21, 44). Recently, it has been demonstrated that CNF-1 mediates its effect via both a clathrin-independent endocytic mechanisms and an acidic-dependent membrane translocation step in its delivery of the catalytic domain to the cell cytosol (12). The consequences of this activation is mainly characterized by an intense actin reorganization (3, 7, 18, 27, 47, 48). We have previously shown that CNF-1, when incubated with epithelial cells, decreases the transepithelial migration of PMNL (26). Moreover, we have also demonstrated that CNF-1 can act on PMNL, by inducing a decreased bacteria phagocytosis (27). No information is currently available on the effect of CNF-1 on T lymphocytes and the possible consequences for the epithelial cells of their interaction with such CNF-1-intoxicated T lymphocytes.

In this study, we present evidence that CNF-1 induces T-lymphocyte phenotypic changes by activation of GTP-binding protein Rho. Such CNF-1-treated T lymphocytes cause cytotoxic effects in cultured monolayers of the human intestinal...
epithelial cell line T84 by increasing their adherence at the basolateral pole of the monolayers. Altogether, these data suggest that during infection by certain E. coli strains producing CNF-1, the adherence and activation status of T lymphocytes may be modified in vivo.

MATERIALS AND METHODS

Reagents. Highly purified CNF-1 used throughout this work was a generous gift of Gilles Flahaut (INSERM U 452, Faculty of Medicine, University of Nice Sophia-Antipolis) and was prepared as described previously (15).

T-cell purification and cell culture. Heparinized blood samples were separated by Ficoll gradient technique. Peripheral blood mononuclear cells were removed from the gradient, and purified T cells were obtained by positive selection as described previously (40). The cells were washed and maintained in RPMI 1640 medium until utilization. The human leukemic T-cell line Jurkat was cultured in RPMI 1640 medium supplemented with 10% (vol/vol) fetal bovine serum, penicillin (100 U/ml), streptomycin (100 μg/ml), and amphotericin B (5 μg/ml). HEp-2 cells were grown in Dulbecco’s modified eagle’s medium (DMEM) supplemented with 7% calf serum, 1% glutamine as previously described (12).

Response to CNF-1 assay. CNF-1-induced Rho activation in T lymphocytes was determined as previously described (21). Deamidation of Ghb by CNF-1 results in a significant upshift of the apparent molecular weight of [32P]ADP-ribosylated Rho on SDS-PAGE (21, 44). Molecular effects due to CNF-1 which result in Rho activation can be demonstrated by radiolabeled staining of Rho and deamidated Rho by the use of C3 exoenzyme, which add [32P]ADP ribose on Thr37/38 of Rho. After sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE), the deamidated protein is detected at a slower migrating band (upshift). Peripheral blood T lymphocytes and Jurkat and HEp-2 cells (10⁷ cells per condition), were incubated with increasing concentrations of CNF-1 (from 1 pM to 3 nM) at 4 and 24 h in supplemented RPMI 1640. Cells were then prepared as previously described (9, 21, 44).

Morphological study. i) Confocal microscopy studies. F-actin fluorescence staining of control and CNF-1-treated cells (peripheral blood T lymphocytes and Jurkat and HEp-2 cells) was processed as previously described (26). The slides were observed and photographed with a laser scanning fluorescence microscope (Leica, DMIRBE, Lyon, France) equipped for epifluorescence.

(ii) Electron microscopy studies. Control and CNF-1-treated cells (T lymphocytes and Jurkat cells) were fixed with 2% formaldehyde in 0.1 M Na cacodylate, pH 7.4, for 1 h at 4°C. Cell pellets were then treated as previously described (26). For immunoelectron microscopy, control or CNF-1-stimulated T lymphocyte pellets were fixed in 3.7% paraformaldehyde and embedded at low temperature into LR White resin (Hard LR, White, London, United Kingdom). Ultrathin sections were put on 300 mesh nickel grids, washed with phosphate-buffered saline (PBS), then incubated for 1 h at room temperature with anti-C11a (Immunotech, Luminy, France), anti-CD29 (K20, gift of M. Tichioni, INSERM 343, Nice, France) or anti-CD103 (β3) (Fib504; American Type Culture Collection [ATCC], Manassas, Va.) antibodies. After being washed with PBS, the grids were incubated for 1 h with 10 nm-thick colloidal gold-conjugated rabbit anti-mouse or goat anti-rabbit secondary antibodies (dilution 1:20; British Biocell International, national, Paris, France). Secondary antibody alone and KL1 (anticytokeratine) antibody (Immunotech) were used in each experiments as negative controls. The grids were washed with PBS, then with distilled water and stained with uranyl acetate. Ultrathin sections were examined with a JEOL 1200 XII electron microscope.

In some experiments, Jurkat cells were incubated with 10⁻³ M CNF-1 for 48 h and then fixed in methanol and stained with May Grumwald-Giemsa (E. Merck, Darmstadt, Germany). To quantify toxic effects in Jurkat cell cultures, 20 microscopic fields (magnification, ×400) for each sample were examined; viable cells (identified by trypan blue exclusion), and cells containing two or more nuclei were counted. Data were expressed as cells per square millimeter.

Flow cytometric analysis. (i) Determination of F-actin. Control and CNF-1-treated lymphocytes were fixed with 3.7% formaldehyde and incubated for 45 min with PBS containing 500 nM rhodamine-phalloidin. After being washed in PBS, the cellular content in F-actin was measured by flow cytometry.

(ii) Expression of integrins. Control and CNF-1-treated lymphocytes were fixed in 1% formalin for 30 min at room temperature. The cells were then washed in Hanks balanced salt solution (HBSS) and incubated with goat immunoglobulin (IgG) (anti-TGF-β) (Peprotech) for 30 min at room temperature. The cells were washed again in HBSS and incubated with either monoclonal antibodies (MABS) (anti-Cd11a; ATCC: diluted 1/1,000), anti-CD29 (K20; INSERM 343; diluted 1/1,000), anti-CD103 (β3) (Fib504; ATCC), or an isotype-match control for 20 min at room temperature and then washed twice. Cells were then treated as previously described (27).

Migration assays. Assays for migration of control and CNF-1-treated T lymphocytes were performed in Transwell cell culture chambers with polycarbonate filters (surface, 0.33 cm²; pore size, 8 μm; Costar). Control T lymphocytes (2 × 10⁵) and CNF-1 pretreated T lymphocytes for 24 h were added to the upper reservoir. One milliliter of HBSS with or without 10 nM SDF-1α (R&D Systems, Lille, France) were added in the lower reservoir. After 6 h of migration, transmigrated cells in the lower reservoir were counted by flow cytometry using a standard of 25 × 10⁴ fluorescent beads (Fluospheres; Beckman Coulter, Paris, France).

Adherence assay to intestinal epithelial cells. Assays for adherence of control and CNF-1-treated Jurkat cells were performed in Transwell cell culture chambers with polycarbonate filters (surface, 0.33 cm²; pore size, 8 μm; Costar). The epithelial intestinal cell line (T84) was grown on these filters as previously described (25, 36). Briefly, inverted monolayers were grown as follows: 0.8-mm-thick lexan rings having the same dimension of the base of Costar inserts were attached to the underside of the insert using general Electric RTV silicone glue. After drying overnight, the inserts were sterilized by submersion in 70% ethanol, inverted onto a sterile petri dish in a hood. Collagen (from rat tail tendon) and T84 cells were added to the filter (underside facing up) and cells were allowed to attach overnight before righting the inserts into the 24-well holding plates. Adherence assays were then performed in a mucosa-to-lumen direction (i.e., Jurkat cells or T lymphocytes were added to the upper reservoir in contact with the basal side of T84 monolayers). Jurkat cells were labeled for 16 h with [3H]labeled thymidine (2.5 μCi/ml; ICN) and preincubated or not at 37°C with CNF-1. A total of 2 × 10⁵ cells suspended in 100 μl of RPMI 1640-0.05% bovine serum albumin was added to the upper chamber and 1 ml of the same medium was added to the lower chamber. When indicated, 10 nM SDF-1α was added to the lower chamber. Cells were allowed to adhere for 6 h at 37°C and 5% CO₂ atmosphere. Counting of adherent Jurkat cells was evaluated by the measure of incorporated [3H]labeled thymidine by scintillation spectroscopy. Data are pooled from 6 to 12 individual monolayers for each condition, and results are means ± standard errors (SE) of five experiments.

Determination of T-lymphocyte adherence to T84 monolayers was performed with a fluorescence multwell plate reader (CytoFlour Perspective Biosystems, Framingham, Mass.). Briefly, 2 × 10⁵ T lymphocytes were added to the upper chamber and 1 ml of HBSS with or without 10 nM SDF-1α was added in the lower reservoir. Cells were allowed to adhere for 6 h at 37°C and 5% CO₂ atmosphere. After 1 h of incubation at 37°C and washing with HBSS, cell-associated fluorescence was measured. Assays were realized in triplicate. The results were expressed as relative mean adherence corresponding to the ratio of fluorescence values before and after washing. In some experiments, adherence of CNF-1-treated T lymphocytes to T84 monolayers that were pretreated with 25 μM pepstatin, an inhibitor of neutral protease activity, was studied in T lymphocytes treated with anti-tumor necrosis factor alpha (anti-TNF-α) MAb (5 μg/ml; Euromedex, Paris, France) or anti-transforming growth factor β1 (anti-TGF-β1) MAb (5 μg/ml; Euromedex) for 1 h.

The morphological consequences of this control T-lymphocyte adherence on T84 monolayers were assessed by electron microscopy. Three different T84 monolayers were examined for each condition after 24 h of contact with lymphocytes treated or not with CNF-1 (16 h, 10⁻³ M). After removal from the inverted, the monolayers were fixed with 2% freshly prepared formaldehyde, in 0.1 M Na cacodylate, pH 7.4, for 1 h at 4°C. Tissues were rinsed in cacodylate buffer and were then processed as described above. More than 30 T84 cells were examined for each condition.

Analysis of activation of p42-44 MAPK and JNK by Western blotting. Control and CNF-1-treated T lymphocytes were washed in Hanks’ balanced salt solution (Sigma, Paris, France), then lysed in cold NP-40 buffer (50 mM Tris-HCl [pH 7.4], 150 mM NaCl, 2 mM EDTA, 1% NP-40, 1 mM Na₃VO₄, 1 mM phenylmethylsulfonyl fluoride, 25 μM leupeptin, 5 mM benzamidine, 1 mM pepstatin, 25 μM aprotinin, 50 mM sodium β-glycosidase, 20 mM sodium pyrophosphate, 0.5 mM dithiothreitol) at a density of 5 × 10⁹ cells/ml. After sonication (two pulses of 8 s), lysates were centrifuged at 15,000 × g for 15 min at 4°C and denatured by boiling in reduced SDS sample buffer. Protein lysates (50 μg per sample) were resolved onto SDS-PAGE and subsequently were electrochemically transferred to polyvinylidene fluoride membrane (Immobilon-P; Millipore). Membranes were incubated in saturation buffer, and probed with anti-phospho-p42-44 MAPK (diluted at 1/2,000) (New England BioLabs, Beverly, Mass.), anti-phospho-JNK (diluted at 1/3,000; New England BioLabs), anti-ERK2 (0.1 μg/ml; Santa Cruz Biotechnologies, Santa Cruz, Calif.), anti-JNK1 (0.1 μg/ml; Santa Cruz,
multiprobe template set from Pharmingen (San Diego, Calif.) (24). Briefly, riboprobes were 32P labeled and hybridized overnight in solution with 1 to 2 pg/ml for TGF-1, TGF-3, or TNF-1, TGF-2, IL-2, IL-4, or IL-8 cytokines, were determined by extrapolation from cytokine standard curves. The minimum labeling the corresponding concentration. Cytokine concentrations in each sample were determined by statistical significance for adherence assays and ELISA analyses. Values are expressed as the means ± SE of the means of a number of experiments.

RESULTS

CNF-1 induces T-lymphocyte phenotypic changes by activation of GTP-binding protein Rho. Since the effect of CNF-1 on lymphoid cells was unknown, we assessed whether T lymphocytes were sensitive to CNF-1. We studied the CNF-1 activation of Rho at different times and concentrations using the Jurkat T-cell line. We used HEP-2 as a CNF-1 sensitive cell line. Rho activation was elicited in a dose- and time-dependent manner (Fig. 1A and B). We then tested this toxin on purified T lymphocytes. We observed the same upshift for T-lymphocyte cells after incubation with CNF-1 (3 \times 10^{-9} M, for 4 h) (Fig. 1C).

The morphology of control and CNF-1-treated T lymphocytes was assessed by electron microscopy. Changes in the shape of CNF-1 stimulated lymphocytes were detectable after 24 h of treatment. Cells exhibited cytoplasmic projections resembling pseudopodia or filopodia. These effects were detected both in peripheral blood T lymphocytes (Fig. 2A and B) and in Jurkat cells (Fig. 2C and D).

Owing to the fact that CNF-1 induces an important F-actin reorganization in various models, we tested the effect of this toxin (10^{-9} M for 24 h) on the T-lymphocyte actin cytoskeleton. The F-actin distribution was investigated by conventional fluorescence and by confocal microscopy after rhodamine-phalloidin staining. In most control cells, an evenly distributed subcortical actin band was consistently observed (Fig. 2E). In contrast, CNF-1 treatment caused the concentration of subcortical actin in broad membrane extension (Fig. 2F). As assessed by flow cytometry, the content of F-actin was increased in CNF-1-treated T lymphocytes (10^{-9} M for 24 h) in comparison to control T lymphocytes (225 ± 11 versus 85 ± 8, means ± SE, for treated cells versus control cells, respectively; P < 0.05) (Fig. 2H and G).

In Jurkat cells incubated 48 h with 10^{-9} M CNF-1, formation
of giant multinucleated cells was found. More than 4 nuclei per cell were observed in 80% ± 10% of the treated population whereas only 5% ± 3% of the control population had more than four nuclei per cell (P < 0.001; not shown).

**CNF-1 induces clustering of CD29 and CD11a integrins on human T lymphocytes.** Quantification performed by flow cytometry showed that exposure of T lymphocytes to CNF-1 for 24 h failed to modify expression of CD11a (Fig. 3A and B) and CD29 (Fig. 3C and D), as well as CD49d and CD103 (not shown). As shown by electron microscopy CD11a and CD29 preferentially clustered into filopodia in CNF-1-treated cells (Fig. 3F and H) while they were evenly distributed along the plasma membrane in control cells (Fig. 3E and G). Identical results were obtained for the CD49d molecule (data not shown).

**CNF-1 potentiates SDF-1α-induced transmigration of T lymphocytes across acellular filters and enhances the adherence of lymphoid cells to T84 monolayers.** As shown in Fig. 4A, pretreatment of T lymphocytes with CNF-1 enhanced SDF-1α induced migration across acellular filters compared with untreated T lymphocytes.

Incubation of Jurkat cells with 10⁻⁹ M CNF-1 for 24 h in the absence of SDF-1α increased four times the number of adherent cells associated with the colonic epithelial cells ([8 ± 1.21] × 10⁴ versus [2 ± 0.9] × 10⁴ cell equivalent, means ± SE, for CNF-1-treated and control cells, respectively; P < 0.05) (Fig. 4B). In parallel, a pretreatment of Jurkat cells with CNF-1 (10⁻⁹ M for 24 h) increased twice their adherence to epithelial cells in response to SDF-1α ([32 ± 1.9] × 10⁴ versus [15 ± 0.9] × 10⁴ cell equivalent, means ± SE, for CNF-1-treated and control cells, respectively; P < 0.05). As shown in Fig. 4A, pretreatment of T lymphocytes with CNF-1 enhanced SDF-1α induced migration across acellular filters compared with untreated T lymphocytes.

**FIG. 2. Morphological modifications and F-actin reorganization induced by CNF-1 (10⁻⁹ M, 24 h) in lymphoid cells.** Transmission electron microscopic photographs of control (A) and CNF-1-treated (B) Jurkat T cells and of control (C) and CNF-1-treated (D) peripheral blood T lymphocytes (magnification, ×2,500). F-actin distribution stained with rhodamine-phalloidin (500 nM) in control (E) and CNF-1-treated (F) peripheral blood T lymphocytes were observed by confocal microscopy (magnification, ×2,000). Determination of lymphocyte F-actin by flow cytometry incubated with CNF-1 (H) exhibited a shift in the fluorescence peak, indicating polymerization of actin compared to control cells (G). One of five experiments is shown (each condition performed in triplicate). P < 0.05.

**FIG. 3. Effect of CNF-1 on CD29 and CD11a expression.** CNF-1 did not cause an increase in CD11a expression (B) or in CD29 expression (D) as assessed by flow cytometry, in comparison with control peripheral blood T lymphocytes (A and C). For CD11a and CD29 staining, numerous beads were regrouped on filopodia in CNF-1-treated cells (anti-CD11a MAb [F]; anti-CD29 MAb [H]) (arrows). No beads were observed in nonfilopodial plasma membrane (arrowheads). Beads were evenly distributed along the plasma membrane in control cells (anti-CD11a MAb [E]; anti-CD29 MAb [G]) (electron microscopy magnification, ×4,500).
1.2] \times 10^4 \text{ cell equivalent, means } \pm \text{ SE, for CNF-1-treated versus control cells, respectively; } P < 0.05 \) (Fig. 4B). We found that CNF-1 enhanced the adherence of T lymphocytes to epithelial cells both in the absence or in the presence of SDF-1α (Fig. 4C).

Analysis of epithelial cells following contact (24 h) with CNF-1-treated lymphocytes by electron microscopy revealed a cytoplasmic vacuolation of T84 cells and a profound alteration of the brush border (Fig. 5A and B) and tight junction disruptions (not shown). By contrast, T84 cells cocultured with un-
treated T lymphocytes showed that the majority of epithelial cells exhibited a regular brush border without disorganization of the monolayers after 24 h (Fig. 5C).

**CNF-1 induces p42-44^{MAPK} activation in T lymphocyte and such activation is required for the increased adherence of T lymphocyte.** The effect of CNF-1 on the different MAP kinase pathways in T lymphocytes was assessed by the use of phosphospecific MAP kinase antibodies. Phosphorylation of p42-44^{MAPK} started after 2 h of incubation with CNF-1, reached a maximal at 3 h and decreased by 4 h (Fig. 6). CNF-1 also induced JNK activation in a comparable time-course (Fig. 6). PMA-mediated activation of p42-44^{MAPK} and JNK via the PKC pathway was used as a positive control (Fig. 6). The extent of JNK and p42-44^{MAPK} phosphorylation was weaker in CNF-1-treated cells than in PMA-treated cells. No activation was observed in control T lymphocytes (data not shown).

As shown on Fig. 7, pretreatment of T lymphocytes with the MEK inhibitor PD 98059 markedly reduced the increase in adherence of CNF-1-treated T lymphocytes to T84 epithelial cells (Fig. 7). Both spontaneous adherence of CNF-1-treated T lymphocytes to T84 cells and adherence of CNF-1-treated T lymphocytes to T84 cells induced in response to SDF-1α were decreased following pretreatment of T lymphocytes with PD 98059 (Fig. 7).

**CNF-1 induced production of TNF-α and TGF-β by T lymphocytes.** Freshly isolated T lymphocytes constitutively produced low levels of cytokines (Table 1 and Fig. 8). CNF-1 treatment of T lymphocytes induced the production of high levels of TGF-β1, TGF-β2, TGF-β3, and TNF-α proteins (Table 1). TGF-β1, TGF-β2, TGF-β3, and TNF-α mRNA were increased in T lymphocytes following CNF-1 treatment (Fig. 8). Incubation of T lymphocytes with CNF-1 had no effect on IL-2, IL-4, IL-8, and IFN-γ mRNA expression and cytokine production (Table 1 and Fig. 8). The results indicated in Table 1 are means ± SE of three different experiments.

p42-44^{MAPK} activation in CNF-1-treated T lymphocytes (for 3 h, 1 nM) was analyzed by Western blotting after preincubation with TNF-α or TGF-β1 MAbs for 1 h. As shown in Fig.

![Image](http://iai.asm.org/) on September 22, 2017 by guest

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**TABLE 1. Cytokine secretion by T lymphocytes following incubation with CNF-1 (24 h, 1 nM)**

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<th>Cytokines</th>
<th>Mean cytokine secretion ± SE (ng/ml)</th>
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<tr>
<td></td>
<td>CNF-1 treated</td>
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<tr>
<td>TNF-α</td>
<td>2.65 ± 0.41</td>
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<tr>
<td>TGF-β1</td>
<td>3.89 ± 0.33</td>
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<tr>
<td>TGF-β2</td>
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<td>TGF-β3</td>
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<td>IFN-γ</td>
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<td>IL-2</td>
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<td>IL-4</td>
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<td>IL-8</td>
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*Results are based on three different experiments. *, P < 0.01.
DISCUSSION

In this report, we demonstrate that T lymphocytes are sensitive to CNF-1. The toxin has been shown to catalyze deamination of the GTP-binding Rho protein, thus inducing an hyperpolymerization of the actin cytoskeleton. Following prolonged incubation with CNF-1, a significant number of Jurkat T cells appeared multinucleated. This is likely due to an alteration in cytokinesis consecutive to the actin cytoskeleton remodeling. This is the first report regarding an effect of CNF-1 on lymphoid cells. In fact, most of the above-mentioned effect of the toxin have been observed with other cell types such as epithelial, endothelial, monocytic and granulocytic cells (7, 18, 27, 47).

Furthermore, we showed that SDF-1α induced-migration of CNF-1-treated T lymphocytes across acellular filters was increased compared to untreated T lymphocytes. Laudanna et al. have reported that Rho is involved in the signaling of chemoattractant receptors to trigger adhesion of leukocytes (32). Moreover, Rho GTP-binding proteins play a crucial role in coupling G protein-linked chemoattractant receptors to integrin-mediated adhesion in leukocytes. Here we demonstrate that adherence of CNF-1-treated T lymphocytes (Jurkat cells and peripheral blood T lymphocytes) to the basolateral pole of intestinal epithelial cells was increased via the Rho GTPases activation. Taken together, these data support the hypothesis that, in vivo, CNF-1 could facilitate the migration of T lymphocytes toward the subepithelial space.

The effect of CNF-1 on the increased adherence of T lymphocytes may be due to (i) Rho activation and a subsequent redistribution at the cell surface of adhesion molecules such as CD11a, CD29, and CD49d, and (ii) p42-44MAPK activation. Previous studies have shown that the effects of Rho are mediated by the downstream kinases, Dial and the Rho-dependent kinase ROCK, probably by increasing myosin contractility, leading to stress fiber bundling and focal adhesion formation (46). In parallel, several studies have clearly demonstrated the involvement of p42-44MAPK in adhesion-mediated signaling (41–43). Fincham et al. have recently shown that active p42-44MAPK was present in cellular adhesion sites (17). In the present study, we report that p42-44MAPK was activated in CNF-1-treated T lymphocytes. Moreover, we could demonstrate that this activation was crucial for the enhanced adherence of T lymphocytes to epithelial cells, since the MEK inhibitor PD98059 prevented the toxin-induced increase adherence.

After their recruitment from the circulation, T cells are present into lamina propria or are intimately associated with epithelial surfaces (2, 37, 45). The latter cells, namely, intraepithelial lymphocytes, constitute a distinct population of T cells that may arise from both thymus-dependent and independent pathways (23, 31, 35, 37). The role of T lymphocytes during the onset of bacterial infection has been poorly investigated. However, it has been demonstrated that T lympho-

FIG. 8. Expression of IL-2, IL-4, IL-8, IFN-γ, TNF-α, TGF-β1 TGF-β2, and TGF-β3 mRNA in CNF-1-treated peripheral T lymphocytes (24 h, 1 nM). RNA expression was analyzed by RPA as described in Materials and Methods. *, P < 0.01.

FIG. 9. (A) p42-44MAPK and JNK activation in CNF-1-treated T lymphocytes are linked neither to TNF-α nor to TGF-β1 production. p42-44MAPK and JNK activation in CNF-1-treated T lymphocytes was analyzed by Western blotting in cells preincubated with anti-TNF-α (lane 3) or anti-TGF-β1 (lane 4) MAbs. T lymphocytes were preincubated for 1 h at 37°C with either anti-TNF-α or anti-TGF-β1 antibodies before addition of 1 nM CNF-1 for 3 h. Lane 1, untreated cells; lane 2, CNF-1-treated cells (3 h, 1 nM); lane 5, PMA-treated cells. (B) Increased adherence of CNF-1-treated T lymphocytes to T84 monolayers is not modified in T lymphocytes pretreated with anti-TNF-α or with anti-TGF-β1 MAbs. T lymphocytes associated with monolayers were counted after 6 h of transmigration (basolateral-to-apical direction) induced by SDF-1α as described in Materials and Methods. Data are pooled from 6 to 12 individual monolayers for each condition, and results are means ± SE (error bars) of five different experiments.
cytes can be activated during the first 24 h of bacterial infection (38, 39). During their activation, T lymphocytes can undergo extensive divisions and acquire effector functions (38). Furthermore, when stimulated, intestinal mucosal T lymphocytes produce various inflammatory cytokines which can elicit different effects on intestinal functions (10, 11, 39). Interestingly, these cytokines can act on PMNL during the acute phase of inflammation induced by bacteria. In this line, IFN-γ was shown to modulate the PMNL migration across T84 cells (10).

Other cytokines produced by stimulated T lymphocytes such as TNF-α can act on PMNL during the acute phase of inflammation induced by bacteria. In this line, IFN-γ can act on PMNL during the acute phase of inflammation induced by bacteria. In this line, IFN-γ can act on PMNL during the acute phase of inflammation induced by bacteria. In this line, IFN-γ can act on PMNL during the acute phase of inflammation induced by bacteria. In this line, IFN-γ can act on PMNL during the acute phase of inflammation induced by bacteria.

In conclusion, we propose that during acute colitis due to certain E. coli strains, CNF-1 toxin could act on T lymphocytes by increasing their adherence to the intestinal epithelial cells. Moreover, by enhancing the production of TNF-α and TGF-β by T lymphocytes, CNF-1 may in turn amplify the inflammatory consequences of PMNL transepithelial migration in response to the bacterial aggression.

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REFERENCES


