Isolation of intestinal mononuclear cells: factors released which affect lymphocyte viability and function

P R GIBSON, A HERMANOWICZ, H J J VERHAAR, D J P FERGUSON, A LÓPEZ BERNAL, AND D P JEWELL

From the Gastroenterology Unit, Radcliffe Infirmary, the Department of Electron Microscopy, John Radcliffe Hospital, and the Nuffield Department of Obstetrics and Gynaecology, John Radcliffe Hospital, Oxford

SUMMARY Whether toxic or immunomodulatory factors are released during the collagenase digestion phase of the isolation of mononuclear cells from human intestinal mucosa was investigated by assessing the effect of the collagenase supernatant on the viability and natural killer activity of normal peripheral blood mononuclear cells. Three hours' incubation in collagenase supernatant suppressed natural killer activity by 25±4% and decreased the viability of peripheral blood mononuclear cells by 11±2%. The ability of collagenase supernatants to kill 51Cr-labelled peripheral blood mononuclear cells over four hours was assessed in 16 collagenase supernatants, eight of which produced lysis of 20±4%. There was no ultrastructural evidence of early degenerative changes in the viable intestinal mononuclear cells fresh from the isolation process or in peripheral blood mononuclear cells incubated in collagenase supernatant. Because prostaglandins are known to inhibit natural killer activity, PGE was measured in 20 collagenase supernatants by radioimmunoassay and found to be high at 27-5±4-0 ng/ml. Addition of indomethacin to the collagenase medium, however, failed to abolish the inhibitory effect of collagenase supernatant on natural killer activity and did not increase the natural killer activity of isolated intestinal mononuclear cells. The release of cytotoxic and immunomodulatory factors during the isolation of intestinal mononuclear cells indicates the necessity for careful assessment of the potential effects of the isolation process on any function being examined and casts doubt upon the relationship between in vitro findings and in vivo functional capabilities.

The enzymatic (collagenase) method for isolating mononuclear cells from intestinal mucosa has been a major advance in the study of local immune mechanisms. It usually yields greater than 2 million cells per gram of mucosa, more than 85-90% of which are viable. Their survival in culture is similar to mononuclear cells isolated from peripheral blood and they are functionally active. The cells proliferate in response to mitogenic stimuli and mediate lectin-induced cellular cytotoxicity, while B cells synthesise immunoglobulins and macrophages show random motility and chemotaxis similar to peripheral blood monocytes.

A major problem in the interpretation of such functional studies has been the disparity of findings between different investigating groups. For example, the proliferative response of intestinal mononuclear cells to phytohaemagglutinin is reported to be less than or equal to that of autologous peripheral blood mononuclear cells. For Crohn's disease, intestinal mononuclear cells from diseased segments have been reported to respond greater than or less than mononuclear cells from histologically normal intestine. Similarly, the pattern of immunoglobulin synthesis differs widely in two reports. Conflicting results have also been obtained using mechanically-isolated intestinal mononuclear cells in suppressor cell assays and in their blastogenic response to mitogens when compared to those using enzymatically-isolated cells. Explanations for these observations include differing methodologies for the functional assays and the heterogeneity within the disease groups studied. The possibility that the functional state of the isolated cells may be variably influenced by the enzymatic extraction process, however, has received little attention. Most investigators have...
confirmed that the EDTA and collagenase used in the isolation procedure have minimal effects on the function of peripheral blood mononuclear cells.3-4 12-14 During the collagenase digestion of the mucosa, however, there is the potential for a release of multiple factors such as proteases and prostaglandins which may profoundly affect the functional status of immunologically active cells.

The aim of this study was to determine if any such immunomodulatory substances are released during the isolation process and the inhibition of natural killer activity was used as a screening test to detect their presence. A factor cytotoxic to peripheral blood mononuclear cells and at least two factors inhibitory to natural killer cells were identified in the supernatant of the collagenase digestion.

Methods

Specimens
Intestinal mononuclear cells were isolated by a modification of the method of Bull and Bookman.1 Intestinal mucosa was obtained from fresh specimens of intestine surgically resected because of Crohn's disease, ulcerative colitis, colorectal carcinoma, and other diseases (colonic adenomata, peptic ulceration of the colon). The degree of inflammation in Crohn's disease and ulcerative colitis specimens is outlined in the results where appropriate. In patients with carcinoma, the tissue examined was taken distant from the tumour and was histologically normal. None of the non-inflammatory bowel disease patients were taking immunologically active drugs whereas all of the Crohn's disease patients and two of four of the ulcerative colitis patients were being treated with high doses of prednisolone (20–64 mg/day). Sulphasalazine had been stopped in all patients before surgery. Heparinised peripheral blood was drawn from normal healthy volunteers.

Isolation of Mononuclear Cells
Within one hour of resection, the intestinal mucosa was dissected and minced in approximately 3 x 3 mm pieces. Six to eight grams of these were incubated with stirring in Hanks balanced salt solution free of calcium and magnesium (HBSS-CMF) with 0.75 mM EDTA and 10 mM HEPES buffer for 45 minutes at 37°C. This was repeated every 30 minutes until no more epithelial cells were seen in the supernatant (usually four to five times). After washing twice with HBSS-CMF, the mucosa now depleted of epithelial cells was incubated with 75–100 ml complete medium (RPMI-1640 with 10% fetal calf serum, 25 mM HEPES, L-glutamine, 100 μg/ml streptomycin, and 100 units/ml penicillin) containing 25 units/ml collagenase (CLSPA, Worthington, Freehold, New Jersey, USA) overnight (approximately 13 hours) at 37°C with stirring. After allowing 20 minutes for the debris to settle, the medium was aspirated and centrifuged at 400 g for 10 minutes to remove remaining debris and cells. The resulting collagenase supernatant was retained and for most experiments used fresh. Aliquots were also stored at -20°C for future prostaglandin assays and for some mononuclear cells preincubation experiments. Storage at -20°C did not affect the potency of collagenase supernatant to inhibit natural killer activity (data not shown). The cell pellet was washed and resuspended in RPMI-1640 and the mononuclear cells purified over a discontinuous Ficoll-Paque (Pharmacia, Uppsala, Sweden) gradient. After washing twice, the mononuclear cells were resuspended in complete medium. Overt microbial infection was excluded by observation of the pH of collagenase supernatant, by microscopy of haematoxylin stained preparations of the isolated cells and by prolonged culture of the intestinal mononuclear cells. The supernatants, however, were not routinely cultured.

Heparinised peripheral blood was diluted 1 in 2 with HBSS-CMF and, within two hours of collection, layered onto a discontinuous Ficoll-Paque gradient and centrifuged at 400 g for 30 minutes.15 The peripheral blood mononuclear cells were harvested, washed twice with HBSS-CMF and suspended in complete medium until used. Mononuclear cells were counted using a haemocytometer (Neubauer chamber) and viability determined using 0.1% trypan blue exclusion.

Cytotoxicity Assays
For the natural killer cell assay, K562 cells were used as the target. 0.5–1x10⁶ K562 cells were incubated in 100 μl of fetal calf serum and 100 μCi of sodium ⁵¹chromate (Amersham, Buckinghamshire, England) for one hour at 37°C. After washing three times, the cells were resuspended in complete medium at a concentration of 1x10⁶ cells/ml. Assays were performed in triplicate or quadruplicate in U-bottomed 96 well microtitre plates (Gibco Bio-Cult, Paisley, Scotland). Fifty microlitres of labelled target cells were added to varying numbers of effector cells according to the effector to target (E:T) ratio to be used. For peripheral blood mononuclear cells, the E:T ratio was 50:1 and for intestinal mononuclear cells 500:1. Each well was made up to a final volume of 200 μl with complete medium. Maximal release was assessed by lysing target cells with Triton-X-100 (BDH Chemicals, Poole, England) in complete
medium and spontaneous release determined by incubating labelled K562 cells in 200 μl of complete medium alone. After four hours' incubation at 37°C and 5% CO₂, 100 μl of cell free supernatant was carefully aspirated and counted in a gamma counter.

For the peripheral blood mononuclear cells cytotoxicity assay, 3×10⁶ freshly-isolated peripheral blood mononuclear cells depleted of platelets by centrifugation for five minutes at 100 g were incubated for one hour in 100 μl of fetal calf serum and 100 μCi sodium ³¹ chromate at 37°C followed by washing three times in complete medium. The cells were then incubated a further 30 minutes in 1 ml fetal calf serum then washed once and resuspended in 2-3 ml complete medium. The assay was also performed in U-bottomed 96 well microtitre plates. Fifty microlitres labelled peripheral blood mononuclear cells were added to 150 μl of collagenase supernatant (experimental release) or complete medium (spontaneous release). Maximal release was assessed as for the natural killer assay. The time of incubation was four hours at 37°C 5% CO₂ and the supernatant was harvested and counted similarly to the natural killer assay. Centrifugation before harvesting was found to be unnecessary.

For all assays, the average of triplicate or quadruplicate wells was taken and the cytotoxicity determined by the following formula: % lysis = (E-S)/(M-S) × 100, where E = experimental release, S = spontaneous release, and M = maximal release. Spontaneous release for K562 cells was 5–10% and for peripheral blood mononuclear cells 7–20% of maximum.

**INDIRECT IMMUNOFLUORESCENCE**

Natural killer cell proportions were measured in peripheral blood mononuclear cells suspensions by indirect immunofluorescence techniques previously described using the monoclonal antibody, anti-Leu-7(HNK-1) (Becton Dickinson, Mountain View, California, USA). Goat-anti-mouse IgM conjugated to tetramethylrhodamine isothiocyanate (Nordic, Tilburg, The Netherlands) was used as the second layer in a concentration of 1 in 10. As control, an IgM antibody to granulocytes was used in place of the first antibody. The resulting smears were examined by fluorescent and phase microscopy and at least 200 mononuclear cells were counted.

**ELECTRON MICROSCOPY**

The cell suspensions were fixed in 4% glutaraldehyde in phosphate buffer pH 7-2 for a minimum of four hours. Pellets were formed by centrifugation and were post fixed in 2% osmium tetroxide for two hours before dehydration in ethanol and embedded in Spurr's low viscosity epoxy. Sections were stained with uranyl acetate and lead citrate before examination in a Philips 301 electron microscope.

The ruthenium red staining was carried out as described by Luft with 0-15% ruthenium red present in both the primary and post fixatives. The thin sections in this case were examined unstained.

**PROSTAGLANDIN ASSAY**

Prostaglandin concentrations in collagenase supernatant were measured by radioimmunoassay. A rabbit antiserum obtained by Dr S P Brennecke (Nuffield Department of Obstetrics and Gynaecology, Oxford) was used. The specificity of this antiserum was tested against a large series of prostaglandins and their metabolites. The only significant cross-reactants were: prostaglandin E₂ (PGE₂) 100%, PGE₁ 100%, PGA₂ 8%, 6-oxo PGE₁ 7%, 13,14-dihydro PGE₁ 6%, PGA₂ 3%, PGE₂a 2%, 13,14-dihydro PGE₂ and PGF₁a 1%. The results were expressed as PGE as the antiserum does not distinguish between PGE₁ and PGE₂.

The assay was performed directly on collagenase supernatant without an extraction step. Samples of collagenase supernatant were diluted 1/20 to 1/80 with potassium phosphate buffered saline (0-1 M, pH 7-4) and incubated in duplicate (0-1 ml) with 0-1 ml antiserum (final dilution 1/2250) and 0-1 ml buffer containing 6 μCi [³H]PGE₂ (SA 160 Ci/mmol; Amersham, Buckinghamshire, England). Tubes containing known amounts of PGE₂ (0-200 pg) were incubated in parallel. After incubation, free and bound prostaglandins were separated by adding 1 ml of a 1-25% charcoal/0-125% dextran 75 suspension and centrifuging at 3000 g for 10 minutes. The radioactivity of the supernatant was measured by liquid scintillation counting. The intra-assay coefficient of variation was 7%. Blanks of buffer alone or complete medium (diluted 1/4 with buffer) were indistinguishable from zero. PGE levels were undetectable in complete medium at the same dilutions as that of the samples of collagenase supernatant.

**STATISTICS**

Data have been expressed as mean ± standard error and were compared using Wilcoxon's rank sum test or Student's t test.

**Results**

**EFFECT OF PREINCUBATION OF PERIPHERAL BLOOD MONONUCLEAR CELLS WITH COLLAGENASE SUPERNATANT ON NATURAL KILLER ACTIVITY**

5–10×10⁶ normal peripheral blood mononuclear cells were incubated in 5–10 ml collagenase supernatant for three hours at 37°C. As a control, a
similar number of peripheral blood mononuclear cells were incubated in parallel in complete medium alone. The cells were then washed twice and resuspended in complete medium and, within three hours, tested for their natural killer activity against K562 target cells. In 35 studies, the natural killer activity decreased from 43±3% to 31±3% without alteration in the proportion of cells with the natural killer (Leu-7+) phenotype (9.0±3.6% compared with 9.1±4.2%). The percentage decrease of cytotoxicity, as presented in Figure 1, was similar for collagenase supernatant from inflamed intestine of patients with Crohn's disease or ulcerative colitis and from histologically normal mucosa. Filtration of collagenase supernatant through 0.22 μm filters (Millipore) did not change the observed suppression of natural killer activity of preincubated peripheral blood mononuclear cells.

**Cytotoxicity of Collagenase Supernatant to Peripheral Blood Mononuclear Cells**

Three hours' incubation of peripheral blood mononuclear cells in collagenase supernatant decreased the total mononuclear cells count from 8.5±1.2×10⁶ to 6.1±0.8×10⁶ and the viability by 11±2% compared with cells incubated in complete medium alone suggesting the presence of a cytotoxic factor (Fig. 2). Supernatants from specimens showing histological inflammation affected viability to the same degree as those from histologically normal intestine (12.3±2.7%, 9.4±2.6%, p>0.1).

Sixteen collagenase supernatants were assessed for their ability to kill ⁵¹Cr-labelled peripheral blood mononuclear cells by incubating the mononuclear cells in an 80% concentration of collagenase supernatant for four hours. Seven produced no cytolysis while, in the other nine, cytotoxicity of 20±4% was seen. In two experiments, collagenase alone in culture medium produced no cytolysis. As detailed in Table 1, the detection of cytotoxins in the collagenase supernatant tended to be associated with a lower than expected intestinal mononuclear cells yield, was more commonly produced from histologically normal tissue (seven of 10) than from inflamed mucosa (two of six), but was not associated with a lower viability of the isolated intestinal mononuclear cells. The ability of collagenase supernatant to inhibit the natural killer activity of preincubated peripheral blood mononuclear cells appeared to be independent of the presence of mononuclear cell cytotoxins (Table 1). The natural killer activity of the isolated intestinal mononuclear cells was tested in 10 of the 12 patients. In all, it was less than 6%, being less than 2% in seven.

The effect of filtering collagenase supernatant through 0.22 μm filters was assessed in seven supernatants exhibiting 25±4% cytotoxicity. Filtering abolished the cytotoxicity in all studies. In two additional experiments, filtration through 5 μm filters also abolished cytotoxicity.

**Electron Microscopy**

The morphology of freshly isolated intestinal mononuclear cells and of normal peripheral blood mononuclear cells incubated in collagenase supernatant for three hours was assessed by transmission electron microscopy. There were numerous mononuclear cells present in both cases as well as a number of degenerative cells and cell
debris which appeared to be phagocytosed by macrophages (Fig. 3). The majority of mononuclear cells appeared viable with normal ultrastructural appearances and exhibited none of the features associated with the early stages of cell death by either apoptosis or coagulative necrosis (Fig. 4). In addition, the physiological intact nature of the plasmalemma was confirmed by the exclusion of the ruthenium red (Fig. 5).

**RELEASE OF PROSTAGLANDINS AND THE EFFECT OF INDOMETHACIN**

The mean concentration of PGE in 20 collagenase supernatants was 27.5±4.0 ng/ml as measured by radioimmunoassay. Significantly more PGE tended to be released from inflamed (38.1±6.3 ng/ml) compared with non-inflamed tissue (19.0±11.9 ng/ml, p<0.05). To inhibit the synthesis of prostaglandins, 10−6 M indomethacin was added to the collagenase-containing medium before the mucosal digestion. This did not affect the intestinal cell yield (data not shown) but did markedly decrease the PGE concentration in the collagenase supernatant (Table 2) to concentrations far below those capable of suppressing natural killer activity in two experiments and to a level at which PGE-mediated inhibition is just detectable in the third. The indomethacin treatment, however, failed to abolish the inhibitory effect of collagenase supernatant on natural killer activity of preincubated peripheral blood mononuclear cells and the natural killer activity of the isolated intestinal mononuclear cells did not change (Table 2).

**Discussion**

This study shows that factors influencing the viability and natural killer activity of peripheral

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**Table 1** The relationship of the cytotoxic and NK-suppressive effects of collagenase supernatant to mucosal histology and to intestinal MNC yield and viability.

<table>
<thead>
<tr>
<th>Patient number</th>
<th>Disease</th>
<th>Tissue</th>
<th>Histology</th>
<th>Intestinal MNC</th>
<th>Collagenase supernatant</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Yield (×10⁶/g)</td>
<td>Viability (%)</td>
</tr>
<tr>
<td>1</td>
<td>Carcinoma</td>
<td>Colon</td>
<td>Normal</td>
<td>2.0</td>
<td>93</td>
</tr>
<tr>
<td>2</td>
<td>Carcinoma</td>
<td>Colon</td>
<td>Normal</td>
<td>2.0</td>
<td>93</td>
</tr>
<tr>
<td>3</td>
<td>Crohn’s disease</td>
<td>Ileum</td>
<td>Inflamed</td>
<td>1.6</td>
<td>90</td>
</tr>
<tr>
<td>4</td>
<td>Crohn’s disease</td>
<td>Ileum</td>
<td>Normal</td>
<td>10.6</td>
<td>93</td>
</tr>
<tr>
<td>5</td>
<td>Crohn’s disease</td>
<td>Ileum</td>
<td>Normal</td>
<td>5.0</td>
<td>92</td>
</tr>
<tr>
<td>6</td>
<td>Crohn’s disease</td>
<td>Colon</td>
<td>Inflamed</td>
<td>10.0</td>
<td>90</td>
</tr>
<tr>
<td>7</td>
<td>Carcinoma</td>
<td>Colon</td>
<td>Inflamed</td>
<td>14.5</td>
<td>85</td>
</tr>
<tr>
<td>8</td>
<td>Carcinoma</td>
<td>Colon</td>
<td>Normal</td>
<td>1.0</td>
<td>85</td>
</tr>
<tr>
<td>9</td>
<td>Carcinoma</td>
<td>Colon</td>
<td>Normal</td>
<td>8.0</td>
<td>91</td>
</tr>
<tr>
<td>10</td>
<td>Carcinoma</td>
<td>Colon</td>
<td>Inflamed</td>
<td>6.2</td>
<td>94</td>
</tr>
<tr>
<td>11</td>
<td>Carcinoma</td>
<td>Colon</td>
<td>Inflamed</td>
<td>4.0</td>
<td>90</td>
</tr>
<tr>
<td>12</td>
<td>Carcinoma</td>
<td>Colon</td>
<td>Inflamed</td>
<td>4.0</td>
<td>95</td>
</tr>
</tbody>
</table>

* Cytotoxicity of 51Cr-labelled PBMNC incubated for four hours in CS.
† Normal PBMNC preincubated in CS for three hours, washed, then tested for NK activity within three hours. Control NK activity was always greater than 20%.
NT = not tested. MNC = mononuclear cells. PBMNC = peripheral blood mononuclear cells.
NK = natural killer. CS = collagenase supernatants.

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**Table 2** Effect of indomethacin on prostaglandin E concentration in and the suppression of NK activity by collagenase supernatants from three patients.

<table>
<thead>
<tr>
<th>Patient</th>
<th>PGE concentration of CS (ng/ml)</th>
<th>NK activity of patients’ intestinal MNC (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>without indomethacin</td>
<td>&lt;2</td>
</tr>
<tr>
<td>2</td>
<td>with indomethacin</td>
<td>25</td>
</tr>
<tr>
<td>3</td>
<td>Incubation of normal PBMNC in CS (change NK activity)*</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>without indomethacin</td>
<td>−33</td>
</tr>
<tr>
<td></td>
<td>with indomethacin†</td>
<td>−39</td>
</tr>
<tr>
<td></td>
<td>NK activity of patients’ intestinal MNC (%)</td>
<td>−20</td>
</tr>
<tr>
<td></td>
<td>without indomethacin</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>with indomethacin†</td>
<td>12</td>
</tr>
</tbody>
</table>

* NK activity of normal PBMNC used in assessing CS from patients 1, 2, and 3 was 35%, 49%, and 18% respectively.
† Experimental values compared with control studies using culture medium containing 10−6 M indomethacin.
Blood mononuclear cells are released during the phase of collagenase digestion in the isolation of mononuclear cells from human intestinal mucosa. Assay of natural killer activity was used to screen for possible immunomodulatory substances for two reasons. Firstly, as the cytolysis of the highly sensitive target cell, K562, occurs rapidly (mostly within one to two hours), the cytotoxicity determined should be a true reflection of the functional status of the natural killer cell fresh from the isolation process. Recovery of the cells during prolonged culture would be minimised. Secondly, the natural killer cell is sensitive to multiple in vitro modulating influences and therefore provides an ideal target for the detection of such factors.

The demonstration that collagenase supernatants are cytotoxic to peripheral blood mononuclear cells is not surprising. Substances causing cytopathic effects to monolayer cell culture are found in intestinal mucosal homogenates and some of these have been partly characterised. The cytotoxic factor in collagenase supernatant, however, was not ultrafiltrable. This suggests the possibility that bacterial contamination of the collagenase supernatant may have been a contributing factor but prolonged cultures and stained preparations of the isolated intestinal cells revealed no evidence for the presence of any bacteria though direct cultures of the collagenase supernatant were not performed. In addition, bacteria were never observed in any sample examined ultrastructurally. Alternatively, particulate matter, of which there are large amounts in collagenase supernatant, may be exerting a nonspecific effect. To support this, filtration through 5 μm filters in two experiments also abolished cytotoxic activity. It is also possible that soluble toxins may be adsorbed onto particulate matter or even adsorbed onto the filter. Thus, the nature of the toxin remains unclear. The collagenase itself is...
not responsible. Cytotoxic factors may be released from dying cells during the isolation but the degree of cytotoxicity exhibited by collagenase supernatant did not correlate with the viability of the isolated intestinal cell population.

The importance of the cytotoxic factor is two-fold. Firstly, it may offer an explanation for unexpectedly low yields of mononuclear cells from inflamed and normal mucosa which occasionally occur. Secondly, it may provide one mechanism by which the functions of mononuclear cells are suppressed. Specific loss of the natural killer (Leu-7+) cell of peripheral blood during preincubation in collagenase supernatant did not occur. No ultrastructural evidence that the viable intestinal mononuclear cells or collagenase supernatant-treated peripheral blood mononuclear cells are damaged could be found but this does not exclude a nonspecific toxic effect on some of their functions. The detection of mononuclear cell cytotoxins, however, bore little relationship to the ability of the collagenase supernatant to inhibit the natural killer activity of preincubated peripheral blood mononuclear cells.

The natural killer suppressive effect of collagenase supernatant acted upon the effector population as in none of the reported experiments was the target cell exposed to collagenase supernatant. Addition of dilutions of collagenase supernatant directly to the natural killer assay also inhibit natural killer activity but the complex cytotoxic and modulating effects on both effector and target cells made interpretation difficult (data not shown).

The factors suppressing natural killer activity appear to be present in the soluble fraction of collagenase supernatant. Previous studies by us14 and others12 13 have shown that collagenase itself is not responsible. Prostaglandins are widely distributed in tissues and are actively produced by many cells including lymphocytes and macrophages. They have multiple effects on immunological functions and prostaglandins of the E series are potent inhibitors of natural killer function.20 26-28 It has been previously shown that prostaglandins are released during the mechanical method of isolating intestinal mononuclear cells and that these influenced the measurement of antibody-dependent cellular cytotoxicity.29 The present study has shown that during the collagenase digestion of intestinal mucosa, concentrations of PGE which have previously been shown to inhibit natural killer activity20 are also produced from both normal as well as inflamed tissue. As might be expected from a recent study,30 greater concentrations of PGE tended to be released from mucosa of patients with active inflammatory bowel disease than from patients without histological inflammation. The role of prostaglandins in suppressing the natural killer activity of freshly isolated intestinal mononuclear cells, however, appears small as inhibition of prostaglandin synthesis with indomethacin failed to increase the natural killer activity of intestinal mononuclear cells. PGE has a short-lived and rapidly-reversible effect on natural killer activity20 28 and a major suppressive contribution would not be anticipated. The indomethacin-treated collagenase supernatant still suppressed natural killer activity though less potently, suggesting that factors additional to prostaglandin release were responsible for the suppression of natural killer activity.

The implication of these findings to the study of natural killer activity of intestinal mononuclear cells is obvious. Previous reports of low or absent activity in isolated mononuclear cells populations4 12 13 31 has been considered to be because of a paucity of natural killer cells within the intestinal mucosa.14 As these cells are exposed during an overnight incubation to factors which potently inhibit peripheral blood natural killer cell function, however, the intestinal natural killer activity being measured may be a gross underestimate.

Of wider importance is the relevance of such factors to the assessment of other immune functions. Whether the effects of collagenase supernatant on natural killer activity are reversible was not determined but the effect of more than 10 hours' incubation of intestinal mononuclear cells in a medium containing such potent toxic and modulating factors must result in the isolation of mononuclear cells with functional capabilities distant from those in vivo. In addition, the influences of collagenase supernatant appear to be unpredictable and independent of underlying mucosal histology. Thus, studies of enzymatically isolated intestinal mononuclear cells are unlikely to yield quantitative information of sufficient accuracy to allow assessment of subtle disease related differences unless such methodological problems can be overcome. This may possibly be achieved by manipulation of the isolation technique or by allowing sufficient time in culture for the cells to recover before subjecting them to in vitro functional assay.

In conclusion, a mononuclear cell cytotoxic, PGE, and other natural killer inhibitory factors are released during the collagenase digestion of human intestinal mucosa. The major implication of these findings is that any in vitro functional assay of intestinal mononuclear cells must be carefully interpreted in the light of the possible artefactual influences the isolation process may have on that
function. They also cast doubt upon the relationship of in vitro intestinal cell function to in vivo activity when enzymatically isolated mononuclear cells are studied.

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