Dendritic cells and scavenger macrophages in chronic inflammatory bowel disease

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SUMMARY  We used enzyme (acid phosphatase [AP]) and immunohistochemical techniques and a set of monoclonal antibodies (CD11, CD5, CD4, CD19, CD8, OKIa), including two recently developed antibodies – for example, HECA-452 (specific for an adhesion molecule on high endothelial venules) and RFD1 (specific for ‘active’ human dendritic cells) to analyse the composition of the gut wall infiltrate of 10 well defined cases of chronic inflammatory bowel disease (CIBD) (six Crohn’s disease (CD), four ulcerative colitis (UC)). Two polar forms in a spectrum of gut mononuclear phagocyte types (CD11+) were identified: at the one extreme scavenger macrophages with blunted projections (AP+, Heca-452-, RFD1-) and at the other extreme, dendritic cells with long dendritic cytoplasmic projections (AP–, Heca-452+, RFD1+). Dendritic cells were mainly found in highly organised lymphoid tissue present at the deeper layers in the gut wall (normal gut: underneath the muscularis mucosae and T-cell areas of lymph follicles [25–30 per follicle]; surrounding the broad zone of scavenger macrophages at the bottom of ulcers (CIBD) and fissures (CD) and in the lymphoid aggregates [25–30 dendritic cells per aggregate] adjacent to granulomas (CD)). These observations can be taken as evidence that exaggerated antigen handling and presentation and stimulation of the immune response takes place at these foci. The observation that scavenger macrophages were localised more superficial, as band like zones (normal gut: subepithelial; mainly surrounding ulcers (CIBD) and fissures (CD)) can be taken as evidence that at these spots the ingestion and degradation of foreign material takes place.

Class II MHC positive dendritic cells are important in the process of antigen presentation to lymphocytes and hence in the initiation of immune responses.1 Dendritic cells were initially described in the T cell areas of lymph nodes and spleen. Some years later it was discovered that skin Langerhans cells also belonged to this group of cells. It is now appreciated that Langerhans cells travel from the skin to the draining lymph nodes through the lymphatics, transporting and exposing skin applied antigens on their surfaces.2–4 A similar antigen presenting cell system of dendritic cells occurs in the normal gut, particularly the mucosa associated lymphoid tissue.5 Dendritic cells have not only been studied in normal immune responses, but also in inflammatory lesions such as rheumatoid arthritis and the granulomas of sarcoid, leprosy, and schistosomiasis. Large numbers of dendritic cells are present in such lesions and their in situ morphological relationship to lymphocytes has extensively been studied.6–10

In situ morphological studies on the presence of dendritic cells in the normal gut wall and that of chronic inflammatory bowel disease (CIBD) focusing on their relationship to other lymphoid cells are scarce.11

The recent development of two monoclonal antibodies which can label dendritic cells is important for such in situ morphological studies: (1) Heca-452, which is specific for a highly glycosylated antigenic
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Determinant (molecular weight 90 kilodalton) of a putative adhesion molecule expressed on high endothelial venules and on dendritic cells, which identifies a unique class II major histocompatibility complex antigen expressed on 'active' dendritic cells present in immune lesions such as rheumatoid arthritis, lepromatous lepra and psoriasis.4–5

The aim of the present study is to analyse the composition of the gut wall infiltrate of 10 patients with CIBD, with special reference to the presence of dendritic cells and scavenger macrophages. In order to identify these cells we used a panel of monoclonal antibodies specific for monocytes/macrophage-like cells including HEC-A-452 and RFD1 in immunohistochemical techniques combined with acid phosphatase enzyme histochemistry.

Methods

Patients

Ten patients with histologically, radiologically, endoscopically, and clinically typical Crohn's disease and ulcerative colitis were studied (Table 1). Six patients with Crohn's disease (four women, two men, mean age 28.5 years, range 20–43). Four underwent an ilealcaecal resection, two a total colectomy. Four patients with ulcerative colitis (two women, two men, mean age 33–2 years, range 27–37). All had had a total colectomy. Controls consisted of four patients (three women, one man, mean age 55–2 years, range 44–77) suffering from gut diseases other than CIBD (Table 1). Two had had a right sided hemicolecction, one a sigmoid resection (all for removal of an adenocarcinoma); one of the patients had a partial ileum resection for removal of a malignant fibrous histiocytoma. Control tissue specimens were taken from normal parts of the bowel unaffected by the malignancies.

Materials

Transmural tissue blocks (colonic as well as ileal) were taken (one block/5 cm) and, frozen in liquid nitrogen and stored at −80°C. Adjacent blocks, were fixed in 4% formalin solution, paraffin embedded, cut and stained (H and E). The surgical specimens fulfilled the macroscopical and microscopical criteria for CD and UC.6–7 Because only typical cases of CD were included all the surgical specimens of patients with CD had fissures and granulomas. Six micrometres cryostat sections (10/frozen tissue block) were cut, air dried (overnight), fixed in acetone (10 min, 21°C) and thereafter stored in a vacuum chamber (−20°C overnight). Sections were stained with an indirect immunoperoxidase technique using the monoclonal antibodies listed in Table 2. Sections were incubated for 60 minutes at room temperature. To block the endogeneous peroxidase the sections were pretreated with 0.0075 U glucoseoxidase (Sigma, St Louis, USA) in 0.1 M D-glucose (Darmstadt, Germany) in PBS for 30 minutes. The second step was a rabbit antimmouse Ig labelled with horse radish peroxidase (Dakopatts, Copenhagen,

Table 1 Patient details

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<tr>
<th>Case</th>
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<td>Total colectomy</td>
<td>CD</td>
</tr>
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<td>F</td>
<td>22</td>
<td>Ileocaecal resection</td>
<td>CD</td>
</tr>
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<td>3</td>
<td>F</td>
<td>43</td>
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<td>UC</td>
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<td>12</td>
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<td>13</td>
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<td>14</td>
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<td>58</td>
<td>Ileal segment resection</td>
<td>MFH</td>
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Table 2 Characteristics of the monoclonal antibodies

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<th>Class differentiation</th>
<th>Antigen</th>
<th>Specificity</th>
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<tr>
<td>RFD1 (M)</td>
<td>–</td>
<td>Class II MHC antigen</td>
<td>‘Active’ dendritic cel B cell subsets</td>
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<tr>
<td>HEC-A-452 (R)</td>
<td>–</td>
<td>gp90</td>
<td>High endothelial venules, dendritic cells</td>
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<tr>
<td>Leu 1 (M)</td>
<td>CD5</td>
<td>T1 (p67)</td>
<td>95% human thymocytes, 95% peripheral T-lymphocytes, most slg + B-CLL cells, no normal lymphocytes</td>
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<tr>
<td>Leu 12 (M)</td>
<td>CD19</td>
<td>B cel (p95)</td>
<td>B lymphocytes</td>
</tr>
<tr>
<td>Leu 3a (M)</td>
<td>CD4</td>
<td>T4 (p55)</td>
<td>Helper/inducer T-subsets</td>
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<td>RIV4 (M)</td>
<td>CD8</td>
<td>T8 (p32-33)</td>
<td>Suppressor/cytotoxic T subsets</td>
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<td>OKIa (M)</td>
<td>–</td>
<td>Class II MHC antigens (p29–p42)</td>
<td>B cells, active T-lymphocytes, macrophages</td>
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<td>FK24 (M)</td>
<td>CD11</td>
<td>C3bi receptor</td>
<td>Monocytes and macrophages</td>
</tr>
</tbody>
</table>

M=monoclonal raised in a mouse; R=monoclonal raised in rat. CD=cluster differentiation proposed at the 'Leeucyte Typing Conference' (Boston, MA, USA, 1984). *Poulter, London, UK; †Duivestijn, Amsterdam, The Netherlands; ‡Becton Dickinson, CA, USA; §Rijksinstituut voor Volksgezondheid en Milieuhygiene, The Netherlands; ¶Ortho, NJ, USA; ¶Monosan/Sanbio, The Netherlands.
Danmark, dilution of 1:25, 30 min). For the rat monoclonal antibody (HECA-452) the second step consisted of a rabbit antirat Ig labelled with horseradish peroxidase (Dakopatts, Copenhagen, Denmark, dilution 40, 30 min). The sections were developed with diaminobenzidine (three min) and counterstained with haematoxylin. A simultaneous detection of HECA-452 and intracytoplasmic acid phosphatase activity was done. The enzymehistochemical staining for acid phosphatase was done according to Burnstone. This acid phosphatase reaction was carried out after the immunohistochemical procedure.

**Results**

On the basis of (a) reactivity with the moab against the putative adhesion molecule identified by HECA-452, (b) reactivity with the moab against the special epitope of Class II MHC determinants identified by RFD1, and (c) the presence of the lysosomal enzyme acid phosphatase, and (d) shape, we were able to identify two large mononuclear phagocyte cells in the normal as well as in the diseased gut wall (Table 3). Both cell types were positive with CD11, a monoclonal antibody specific for C3bi receptors indicating, that both cell types were of mononuclear phagocyte character. The first cell type was a large cell (Table 3), which was strongly positive for the lysosomal enzyme AP. Though some expressed Class II MHC determinants (OK1a) the cells did not react with HECA-452 and RFD1. These cells often showed small blunted projections (Fig. 1A). Hence these cells had the phenotypic characteristics of a scavenger macrophage.

The second cell type (Table 3) showed strong reactivity with the moab OK1a, HECA-452, and RFD1 (Fig. 1B); however, practically no AP-activity was identified. Although the saxe was difficult to judge (because of the use of frozen sections), long dendritic cytoplasmic projection were often identified. This cell type most likely represents the dendritic cell active in the handling and presentation of antigen.

These two mononuclear cell types formed the extremes of a spectrum of CD11 positive cells, which showed enzyme and immunohistochemical characteristics positioned in between these two polar forms: such intermediate cell types showed some forms of cytoplasmic protrusions, all had class II MHC determinants (OK1a), and the majority of these cells showed a weak reactivity with the moab HECA-452 and had some AP-activity. It is important to note that these cells did not react with the moab RFD1 (we further refer to such cells as 'undeterminate monocytoïd cells').

**Table 3: Staining characteristics of the dendritic cell and scavenger macrophage**

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Dendritic cell</th>
<th>Scavenger macrophage</th>
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<tbody>
<tr>
<td>CD11</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>OK1a</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>HECA-452</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>RFD1</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>Enzyme: AP</td>
<td>−</td>
<td>+</td>
</tr>
</tbody>
</table>

**Healthy ileum and colon**

Underneath the surface epithelium of the normal colon clusters of zonally arranged scavenger macrophages were found (Figs. 2 and 3a). These cells intermingled with a few dendritic cells. In the villi of the normal ileum such zonally arranged cells were also present but their numbers were lower (approximately two times lower as compared with colonic mucosa).

Deeper down in the lamina propria of the ileum and the colon a few scattered ‘undeterminate monocytoïd’ cells were found, while just above the muscularis mucosae an occasional dendritic cell could be identified. These cells often formed small clusters in which also a few T-lymphocytes could be identified (Fig. 3a).

At the transition of the lamina propria and submucosa, in the area of the muscularis mucosae, an occasional lymphoid follicle was present (approximately two to three per cm bowel segment). The centre of such a follicle consisted of the well known ‘tingible body’ macrophages and B cells (CD19); this B centre was surrounded by a rim of T cells. In this T cell area dendritic cells could clearly be identified (approximately 25–30 per lymphoid follicle) as well as some high endothelial venules, which were, like the dendritic cells, positive for the determinant recognised by the moab HECA-452 (Fig. 3a). The submucosa itself contained a few solitary scavenger macrophages and some ‘undeterminate monocytoïd cells’, but dendritic were largely absent. In the muscularis propria dendritic cells could not be identified and, T- and B lymphocytes were almost absent.

**CIBD**

The bottom of the ulcers (both in CD and UC) and the fissures (only in CD) consisted of a broad zone of densely packed scavenger macrophages, intermingled with numerous ‘undeterminate monocytoïd cells’ (Fig. 3b). This inner zone of mononuclear phagocyte cells was in turn surrounded by an outer zone of cells predominantly consisting of T lympho-
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Fig 1  (A) Scavenger macrophage with small blunted projections (AP+, Heca-452−); (B) Dendritic cell with cytoplasmic protrusions (AP−, Heca-452+) showing contact with T lymphocytes. Immunoperoxidase double staining for AP and Heca-452.

cytes, with a T helper (Th) to T suppressor/cytotoxic (Ts/c) ratio of 2:1. In between these T cells of the outer zone a few dendritic cells could be identified, and some HECA-452 positive high endothelial venules were part of this area (Fig. 3b). Besides these structures a few B cells (CD19) as well some plasma-cells were found in the outer zone, there was also an occasional a lymphoid follicle.

In CD, the granulomas were localised transmurally, often along the draining lymphatics. The packed epithelioid cells forming the centre of such granulomas showed strong acid phosphatase activity; they did however not react with the moabs HECA-452 and RFD1, which probably means that these cells are mainly involved in processes of degradation. Adjacent to or around these epithelioid cells accumulations of T lymphocytes were present. Several high endothelial venules positively staining with the moab HECA-452 were present in these T cell areas (Figs. 3c, 4). The parts of the areas most closely positioned to the epithelioid cells predominantly consisted of Th cells intermingled with dendritic cells (approximately 25–30 per lymphoid aggregate); Ts/c cells were scarce (Th to Ts/c ratio of 6:1). The outer parts of the lymphocytic zones contained relatively higher numbers of Ts/c cells and Th to Ts/c to ratio's of 3:2–2:2 were found. A few B cells could also be identified in these parts, but plasmacells were absent. There were no remarkable differences in the composition of the infiltrate between ileal and colonic tissue in CD.

In UC, the number of lymphoid follicles at the transition of muscularis mucosae had increased (approximately six to eight follicles per cm bowel segment) compared with CD and normal subjects; however, the composition of the follicles was comparable with that of lymphoid follicles present in the normal colon.

Both in CD and UC, the remaining part of the swollen edematous submucosa contained large numbers of scavenger macrophages and ‘undeterminate monocytoid cells’ (four to five times as much as in healthy colon). These cells did not show any noteworthy contacts with other lymphoid cells present in this area.
Discussion

In this immunohistochemical study on surgical specimens of normal gut and typical cases of CIBD, in which two recently developed monoclonal antibodies - that is, Heca-452 and RFD1 were included, we were able to distinguish a dendritic cell (OK1a+, AP-, Heca-452+, RFD1+), and a scavenger macrophage (OK1a+, AP+, Heca-452-, RFD1-) (Table 3).

In normal gut scavenger macrophages were predominantly found zonally arranged just underneath the enterocytes in both colon and ileum. In CD and UC scavenger macrophages were present in higher numbers, as compared with the normal gut. They were arranged in band like zones forming the bottom of the ulcers or fissures. It is likely that the zonally arranged macrophages represent a scavenging defence mechanism towards microbial agents or dietary substances, penetrating through small (in normal gut) or widespread mucosal defects (in case of CIBD). In the study of Selby et al11 on colonic mucosal biopsies of CIBD scavenger macrophages were identified on the basis of the absence of the enzyme ATP-ase, and on the presence of the enzymes AP and non-specific esterase. Though they found a distribution of scavenger macrophages in the normal gut similar to that found by us, the authors reported decreased numbers of such cells in both CD and UC, which contrasts to our findings. An explanation of this discrepancy might be their use of biopsy material which necessarily only includes on analysis of the infiltrate of the superficial mucosal layer. We used surgical specimens, and therefore were able to analyse the deeper positioned infiltrates, surrounding the ulcers and fissures. In normal gut dendritic cells were mainly found in small clusters just above the muscularis mucosae, or just underneath it as parts of the T cell zones of lymphoid follicles situated at the transition of the lamina propria and the submucosa.

In CIBD, the majority of the dendritic cells were found as part of organised lymphoid cell accumulations, predominantly consisting of T cells. These T cell accumulations surrounded the band like zones of scavenger macrophages forming the bottom of the ulcers (CIBD) and fissures (CD), or were found adjacent to the epithelioid granulomas (CD). It is likely that dendritic cells play a role in antigen presentation to these T cells.

Immune granulomas, similar to those found in CD, have been described in the lesions of tuberculosis leprosy, sarcoid and protozoan infections as schistosomiasis.7-15 The granulomas in these disorders are thought to be the result of an exaggerated, but intact T-cell mediated immune reaction towards a persisting antigen of either unknown origin (sarcoid) or of microbial nature (M leprae, the nematode egg).

It is tempting to speculate that the T cell granulomas in CD are likewise engaged in the handling and/or degradation of a persisting antigen. A microbial antigen which has recently come into the focus of attention is that of M paratuberculosis,20 which organism has been recently cultured from gut tissues of patients with CD.21-22 Other antigens that might play a role are those of commensal microbes - for example, E coli 014, those of dietary origin such as cow's milk proteins, or autoantigens such as colonic epithelial cells.23-25

Selby et al in their study on CIBD identified dendritic cells on the basis of strong ATP-ase activity in combination with a weak AP and non-specific esterase activity.11 In contrast with our results, they found such cells to be present in CIBD in equal numbers as in the normal gut wall. In colonic mucosa of patients with CIBD, however, such cells were...
scattered diffusely throughout the lamina propria. Their use of histochemical criteria and mucosal biopsies may have played a role again in the discrepancy between theirs and our data. Further a strong overlap of 'their dendritic cells' with 'our undeterminate monocytoid cells' is probably also the case. (We found many 'undeterminate monocytoid cells' present in the submucosal tissue.)

In conclusion dendritic cells (CD11+, Heca-452+, RFD1+, AP−) and scavenger macrophages (CD11+, Heca-452−, RFD1−, AP+) could be identified in situ in normal and diseased gut wall. They form the extremes in a spectrum of gut mononuclear phagocytes.

Dendritic cells were predominantly identified in highly organised lymphoid tissue, present at the deeper layers in the gut wall (normal gut: underneath the muscularis mucosae; surrounding the broad zone of scavenger macrophages at the bottom of ulcers (CIBD) and fissures (CD), and adjacent to the granulomas (CD). These observations can be taken as evidence that exaggerated antigen handling and presentation and stimulation of the immune response takes place at these foci. The factual presence of these granulomas in CD and their position adjacent to lymphatics as far as the serosal layer and the mesenteric lymph node indicates that the antigens to be handled in this disorder are extremely difficult to degrade. The observation that scavenger macrophages were localised more superficially, as band like zones (in normal gut: subepithelial; mainly surrounding ulcers (CIBD) and fissures (CD)) can be taken as evidence that at these sites there is an exaggerated exposure to and degradation of foreign material.

The differences in pattern of scavenger macrophages and dendritic cells between normal and diseased gut are mainly quantitative. Functional studies should be undertaken, however, to analyse if...
Fig 4  Composition of a granuloma in CD.  
Epithelioid cells (AP+) in the centre, surrounded by a zone of T lymphocytes with some HEV’s (Heca-452+), intermingled with dendritic cells (AP−, Heca-452+); E=epithelioid cell, →=dendritic cell.  
Immunoperoxidase double staining for AP and Heca-452.

there is no defect in the function of these particular cell types.

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