Quantitative distribution of immunoglobulin-producing cells in gastric mucosa: relation to chronic gastritis and glandular atrophy

K Valnes, P Brandtzaeg, K Elgjo, and R Stave

SUMMARY Immunoglobulin (Ig)-producing immunocytes were quantified by paired immunofluorescence staining in specimens of gastric antral (n=52) and body (n=117) mucosa obtained from 45 patients with various gastrointestinal disorders. Enumerations were carried out in a 500 μm wide zone from the muscularis mucosae to the lumen ('tissue unit'). The specimens were divided into three categories according to the degree of inflammation, and each specimen received a grade for atrophy (0–2). The total number of IgA-, IgM- and IgG-producing cells per tissue unit increased strikingly with increasing degree of inflammation, both in antral and body mucosa. IgA immunocytes predominated (61–91%) in all specimens, but the IgG isotype showed the largest relative increase (four to 17-fold), particularly in the basal part of the mucosa. In this layer of the gastric body the proportion of IgG cells was also significantly raised in association with atrophy, irrespective of degree of inflammation. Locally produced IgG may be of protective significance in terms of internal (or 'second line') defence but may at the same time maintain immunopathological mechanisms contributing to the chronicity of gastritis.

Histopathological characterisation of chronic gastritis is usually based on the density of inflammatory cells and the degree of mucosal atrophy. The immunoglobulin (Ig)-producing plasma cells and their immediate precursors (B-cell blasts) are important local effector cells, both in terms of mucosal protection and immunopathology. This protection involves primarily specific humoral defence by the secretory IgA (and IgM) system; after being produced by local immunocytes, dimeric IgA and pentameric IgM are selectively transported by means of secretory component as a receptor through restricted parts of the gastric epithelium.

It has recently been shown that epithelial expression of secretory component and uptake of IgA are enhanced in gastritis. The associated cellular IgA response in the lamina propria, however, related to the degree of inflammation has not been previously quantified. Also, the relative participation of immunocytes of other isotypes (IgM, IgG, IgD, and IgE) in gastritis has not been conclusively established, whereas such aspects of local immunity have been extensively defined in normal and diseased intestinal mucosa. In the present study, therefore, immunohistochemical quantitation of Ig-producing cells was related to chronic gastritis and glandular atrophy in human stomach mucosa.

Methods

SPECIMENS AND TISSUE PREPARATION Specimens of gastric antral (n=52) and body (n=117) mucosa were obtained from 21 women and 24 men with a median age of 43 years (range, 15–72 years). The material was collected surgically from 21 individuals of the following categories: patients subjected to Billroth II resection for duodenal or gastric ulcer (n=15); patients operated for duodenal or gastric neoplasia (n=2); and kidney donors (n=4). In addition, biopsy specimens obtained through a Crosby's capsule from patients with coeliac disease (n=11) or 'non-ulcer dyspepsia' (n=13) were included.

The tissue samples were excised from macroscopically normal mucosa or, when taken with the Crosby's capsule, from stomachs without endo-
scopically or radiographically overt lesions. After prewashing in 0-01 M phosphate buffer, pH 7-6, containing 0-15 M NaCl (PBS), the specimens were fixed in cold 96% ethanol and embedded in paraffin.\textsuperscript{11}

**IMMUNOHISTOCHEMICAL REAGENTS AND STAINING PROCEDURES**

Antisera to the five human Ig isotypes were raised in rabbits and rendered monospecific by appropriate solid phase absorptions.\textsuperscript{12,13} The IgG fractions of the antisera were labelled with fluorescein isothiocyanate or tetramethylrhodamine isothiocyanate\textsuperscript{12}; the characteristics of the purified conjugates are shown in Table 1. Performance tests on negative and positive substrates, and absorption controls with appropriate soluble antigens, confirmed the specificity of immunohistochemical staining.\textsuperscript{13,14}

Serial sections were cut at 6 µm perpendicular to the mucosal surface, dewaxed, and stained either with a haematoxylin, azophloxine, and saffron trichrome method\textsuperscript{15} or by direct paired immunofluorescence with various conjugate combinations as shown in Table 1. The major Ig classes (IgA, IgM, and IgG) were thus shown alternately by green and red fluorescence in adjacent sections. Direct paired immunofluorescence stained sections were regularly mounted in polyvinyl alcohol. Representative sections were remounted in polyvinyl alcohol containing para-phenylenediamine to preserve fluorescence for photographic documentation of paired staining features.\textsuperscript{16}

**MICROSCOPY, CELL COUNTING, AND EVALUATION OF RESULTS**

Direct paired immunofluorescence stained preparations were examined in a Leitz Orthoplan microscope equipped with a Ploem-type vertical illuminator.\textsuperscript{17} Double exposed colour slides (Ektachrome ASA 400 daylight film), were taken with an X10 oil immersion objective to show simultaneously green and red cytoplasmic fluorescence. The slides were projected at a final magnification of ×500 to facilitate counting of fluorescent immunocytes with a discernible nucleus and purely green or red cell profiles with a diameter >8 µm.

IgA-, IgM-, and IgG-producing immunocytes were evaluated in three serial sections with the conjugate combinations shown in Table 1. Two mucosal 'tissue units', each comprising a 500 µm wide section area of the mucosa at full height (from the muscularis mucosae to the lumen), were evaluated in every section.\textsuperscript{5} Immunocytes of each of the three major isotypes were thus enumerated in two units with each conjugate. The mucosa was divided in a luminal 200 µm zone and a remaining basal zone. Arithmetic means of the four counts for IgA-, IgM-, or IgG-positive cells were calculated for each specimen. The average percentage isotype distributions within each mucosal zone and for the whole tissue unit were also determined. Results for various categories of specimens were based on the sample means and standard deviations, and comparisons were done by Wilcoxon's test for unpaired samples (two-tailed) with a chosen level of significance at 5%. Kendall's τ test was used for correlation analyses.\textsuperscript{18}

Histopathological features were evaluated blindly in HAS-stained sections by an independent observer. Grading of chronic gastritis was based on the classification proposed by Rao\textsuperscript{19}; both the antral and the body material were thereby divided into three groups (A, B, C) with increasing degree of inflammation. To obtain groups of an acceptable size, this division had to be different for antral and body specimens because the inflammatory changes were much more extensive in the former. Thus, for the pyloric antrum, group A included only one specimen of grade 0 along with 24 of grade 1, whereas groups B and C represented grades 2 and 3, respectively. For the gastric body, group C included only specimens of grade 2, whereas groups A and B represented grades 0 and 1, respectively. In addition each specimen received a score for atrophy (0 to 2). Method reproducibility of cell counting and grading of inflammation have been reported previously.\textsuperscript{4,20}

**Results**

Ig-producing immunocytes of the three main isotypes were clearly visualised in the lamina propria in

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**Table 1: Applied combination and characteristics of rhodamine and fluorescein conjugates used for quantitation of immunoglobulin-producing cells by paired immunofluorescence staining**

<table>
<thead>
<tr>
<th>Code and labels of combined conjugates</th>
<th>Isotype specificity of conjugates</th>
<th>Optical density (OD) ratio*</th>
<th>Working conc. † (g IgG/litre)</th>
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<tbody>
<tr>
<td>1 131AB TRITC</td>
<td>α chain (IgA)</td>
<td>3-1</td>
<td>0-20</td>
</tr>
<tr>
<td>2 38B FITC</td>
<td>γ chain (IgG)</td>
<td>2-0</td>
<td>0-46</td>
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<tr>
<td>3 137B TRITC</td>
<td>μ chain (IgM)</td>
<td>4-9</td>
<td>0-50</td>
</tr>
<tr>
<td>4 307AB TRITC</td>
<td>α chain (IgA)</td>
<td>2-2</td>
<td>1-02</td>
</tr>
<tr>
<td>5 158B FITC</td>
<td>ε chain (IgE)</td>
<td>4-8</td>
<td>0-30</td>
</tr>
</tbody>
</table>

*OD\textsubscript{290nm}/OD\textsubscript{max} as an estimate of degree of labelling.\textsuperscript{13} † For 20-h incubation at room temperature.\textsuperscript{14} TRITC=tetramethylrhodamine isothiocyanate. FITC=fluorescein isothiocyanate.
Immunocytes in gastric mucosa

Fig. 1  Localisation of IgA-, IgG- and IgM-producing immunocytes in gastric antral mucosa. (a) Section showing paired staining for IgA (left panel, red fluorescence) and IgG (right panel, green fluorescence). (b) Section below shows paired staining for IgA (left panel, green fluorescence) and IgM (right panel, red fluorescence). Lumen at the top. Bar=50 μm.
Valnes, Brandzaeg, Elgjo, and Stave

the prewashed, ethanol-fixed specimens (Fig. 1). IgD- and IgE-producing cells amounted to much less than 1% and could thus be discounted. Mast cells with peripheral IgE were often seen but were not considered in this context.

Some of the diffusely distributed extracellular IgG and IgA were occasionally retained due to incomplete extraction during the prefixed washing process. In double exposed colour slides of direct paired immunofluorescence-stained sections, however, such background staining appeared yellow, thereby facilitating counting of the purely green or red immunocytes.

**Relation to Degree of Gastritis**

*Quantitation of Ig-producing cells in antral mucosa*

Both totally and within each mucosal zone, the number of IgA and IgG immunocytes rose significantly with increasing degree of inflammation, whereas for IgM cells significant differences were found only between groups A and B and between groups A and C (Fig. 2a). From group A to group C the total immunocyte number per mucosal tissue unit increased from 54 to 260 for IgA, from 8 to 25 for IgM, and from 11 to 97 for IgG (Fig. 2a). The highest relative increase was seen in the basal zone for IgG cells, which rose nine-fold from group A to C.

*Isotype proportions of Ig-producing cells in antral mucosa*

IgA immunocytes showed a striking predominance. In inflammatory group A they accounted for 89% and 78% of the Ig-producing cells in the luminal and basal zone, respectively (Fig. 3a). The proportion of IgA cells did not vary significantly between groups in the luminal zone, whereas in the basal zone (and totally) significantly lower proportions of IgA cells

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![Diagram](http://gut.bmj.com/10.1136/gut.27.5.505)

Fig. 2. Number of cells with cytoplasmic IgA, IgM, or IgG in tissue specimens of various inflammatory groups (see text) from gastric antral (A, n=25; B, n=19; C, n=8) or body (A, n=22; B, n=72; C, n=23) mucosa. Upper and lower parts of columns represent luminal and basal zones (see text) of mucosa, respectively. Result given as mean and standard deviation (vertical bar). Significantly increased cell counts compared with one (*) or both (**) of the other groups are indicated (for total mucosal unit, above the column; for different zones, within the respective columns). n=number of specimens.
Immunocytes in gastric mucosa were found in inflammatory groups B and C (61% and 62%, respectively) compared with group A (78%). IgG cells increased significantly with increasing degree of gastritis (luminal zone, group C > A and B; basal zone and totally, groups B and C > A) (Fig. 3a). For all inflammatory groups the percentage of IgA cells was higher in the luminal zone than basally contrary to that of IgG cells (Fig. 3a). The proportion of IgM immunocytes did not vary significantly between the different inflammatory groups or mucosal zones; they accounted for 5–10% of all Ig-producing cells.

Quantitation of Ig-producing cells in body mucosa

The total cell count rose for all major immunocyte classes with increasing degree of inflammation (Fig. 2b). The same held true when the luminal and basal zones were analysed separately (Fig. 2b). From group A to C, the total number of immunocytes per mucosal unit increased from 21 to 124 for IgA, from 3 to 24 for IgM, and from 3 to 55 for IgG (Fig. 2b). As in the antral mucosa, the highest relative increase was found for IgG cells in the basal zone, where a 17-fold rise was revealed from group A to group C.

Isotype proportions of Ig-producing cells in body mucosa

IgA immunocytes predominated in all inflammatory groups and in both mucosal zones, accounting for 57–86% of the Ig-producing cells. The proportion of IgG cells increased both luminally and basally in group C compared with the two other groups (Fig. 3b), whereas the representation of IgA cells decreased (Fig. 3b). As for the antral mucosa, the proportion of IgA cells was significantly higher luminally than basally in all groups whereas the reverse was true for IgG cells. No significant

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**Fig. 3** Percentage isotype distribution of cells with cytoplasmic IgA, IgM, or IgG in tissue specimens of various inflammatory groups (see text and Fig. 2) from gastric antral (a) or body (b) mucosa. Result given as mean and standard deviation (vertical bar). For each isotype significant differences of one group compared with both of the others in the two mucosal zones and in the total unit are indicated (*).
differences in the proportion of IgM immunocytes could be found between the various inflammatory groups or mucosal zones; these cells accounted for 8–14% of the total local immunocyte population.

**Relation to glandular atrophy**
The relation between degree of atrophy and gastritis is shown in Fig. 4. For further statistical analysis of Ig-producing cells related to glandular atrophy, antral and body specimens with (n=33 and n=28, respectively) and without (n=19 and n=89, respectively) atrophy were first compared disregarding the degree of inflammation.

**Ig-producing cells in antral mucosa**
The total number of IgA and IgM immunocytes per mucosal unit was 122 and 15, respectively, in specimens with atrophy compared with 85 and nine in those without; however, this apparent increase was not statistically significant. The number of IgG cells was virtually the same in both categories. The same held true with regard to isotype proportions.

**Ig-producing cells in body mucosa**
Compared with specimens without, those with atrophy had a raised number of IgA (from 55 to 103), IgM (from 7 to 19), and IgG (from 13 to 45) immunocytes per mucosal unit. These differences were statistically significant, also in each mucosal zone. When the isotype proportions were considered, that for IgA cells were significantly decreased and that for IgG increased in both zones of specimens with atrophy compared with those without.

**Ig-producing cells in specimens with glandular atrophy and similar degree of inflammation**
As glandular atrophy and inflammatory changes were significantly correlated (Kendall's τ test) in the body mucosa (Fig. 4), specimens with the same degree of gastritis (either grade 1 or 2, Fig. 4) with or without atrophy were compared. In the body mucosa, specimens of grade 1 (group B) had a significantly higher number of IgG cells in the basal zone in the presence than in the absence of atrophy (Fig. 5). This held true also when the isotype proportions were considered, both in the basal zone (Fig. 5) and in the whole mucosal unit. No such differences were found in grade 2 inflammation or in the antral material.
Discussion

IgA-producing cells were found to be the predominant immunocyte class in both normal and inflamed human gastric mucosa as previously shown for all other secretory tissues investigated in this laboratory.\(^1\) Relatively few immunocytes were present in normal or slightly inflamed mucosa, however, whereas a striking numerical rise was demonstrated with increasing degree of gastritis. Although this rise included all three main isotypes, it was relatively largest for IgG-producing cells.

Local production of IgA is the crucial basis for secretory immunity in man;\(^3\) and restricted parts of the gastric epithelium produce SC\(^4\)\(^5\)\(^22\)\(^23\) and are thereby involved in receptor mediated external translocation of IgA.\(^3\)\(^24\) We and others have found that such epithelial transport is apparently enhanced in association with gastritis.\(^4\)\(^5\) The present study showed that, in addition to increased epithelial expression of SC,\(^4\)\(^5\) a raised number of IgA-producing cells contributes to reinforcement of secretory immunity in gastritis.

Studies of other mucosae have indicated that when insufficient antigen exclusion is afforded by the secretory immune system, a second line of defence develops involving mainly local production of IgG.\(^1\) The relatively high number of IgG-producing cells, particularly in the basal layer of inflamed gastric mucosa, may thus be of biological significance in terms of mucosal protection. As IgG antibodies are complement-activating they may not only retard influx of specific antigen but also increase epithelial penetrability for unrelated macromolecules.\(^25\)\(^26\) Such phlogistic mechanisms, resulting from an altered mucosal homeostasis,\(^1\) may contribute to perpetuation of chronic gastritis and development of glandular atrophy. In addition to activating complement, IgG antibodies can arm K cells which may exert cytotoxicity of target cells. Although little as yet is known about the antibody specificities of gastric immune responses, local IgG cells with a cytoplasmic product binding intrinsic factor have been found in pernicious anaemia.\(^27\) Such antibodies may conceivably cause damage of gastric glands. In simple chronic gastritis, serum autoantibodies to gastrin-producing (G) cells have been reported\(^28\) and these may well be of local origin.

When body specimens with the same grade of gastritis were compared in relation to the presence or absence of atrophy (Fig. 5), the total number of IgG-producing cells per mucosal unit was the same in both groups. In specimens with gastritis of grade 1 (group B), however, IgG cells were relatively more increased in the basal zone of the body mucosa in association with atrophy. The lack of such a relation in gastritis of grade 2 might be explained by the higher overall density of IgG cells in this group; the same was true in the antral mucosa where inflammatory grades 1 and 2 signified more severe gastritis than the comparable grades assigned for the body mucosa.

Several difficulties exist in comparing our results with those from previous immunohistochemical studies of IgG-producing cells in human gastric mucosa (Table 2). The gastric region under investigation has not always been specified\(^29\)\(^30\) and histopathological classification of the specimens has often been lacking\(^33\) or based on different criteria.\(^23\)\(^32\) In some papers, quantitative data have not been reported\(^30\)\(^32\) or they have been based on estimation of cell densities\(^23\)\(^29\)\(^30\)\(^32\)\(^33\) rather than on the total number of immunocytes in a mucosal tissue unit as determined in the present study. Generally, when normal and inflamed gastric mucosae have been compared, most studies have shown an absolute increase in the numbers of cells producing one or more of the three main Ig isotypes.\(^29\)\(^31\)\(^34\)\(^37\)\(^38\) Nevertheless, a clear-cut and significant numerical immunocyte increase according to the histopathological grade of gastritis has not been shown in previous studies.

When the immunocyte isotype proportions are considered, four previous studies give data for normal antral mucosa\(^31\)\(^35\)\(^36\)\(^37\) that show a percentage of IgA (range, 63–88%), IgM (range, 4–23%), and IgG (range, 6–21%) cells comparable with our findings in normal and slightly inflamed antral mucosa (Table 2). The same is true in three studies reporting data for normal body mucosa,\(^31\)\(^35\)\(^36\) in which the median values for IgA (76%), IgM (9%), and IgG (13%) cells fit well with ours (Table 2).

Three previous\(^31\)\(^36\)\(^37\) studies on antral gastritis can be compared with ours – that of Slauoi\(^36\) reporting very similar data. Conversely, the IgG figure is smaller in the study of Spreeuw,\(^37\) and the IgM percentage reported by Camilleri\(^31\) is relatively high. Our result for inflamed body mucosa is also most like that of Slauoi\(^36\) whereas Camilleri\(^31\) again reports strikingly high IgM figures (33%).

The normal antral IgA- to IgG-cell ratio ranged in four previous studies from 4 to 14.\(^31\)\(^35\)\(^36\)\(^37\) In three, a decrease in this ratio was found in connection with gastritis,\(^31\)\(^36\)\(^37\) In our study the same ratio decreased from 5 to 3 with increasing grade of antral gastritis.

For normal body mucosa we calculated an IgA-to IgG-cell ratio of 6 which was similar to that reported in three previous studies.\(^31\)\(^35\)\(^36\) A decrease of this ratio with increasing degree of gastritis was shown by us and others with two
exceptions. Such a decrease was reported to be particularly marked in atrophic gastritis associated with pernicious anaemia. Odgers and Wangel described two such patient groups, one with a normal gastric IgA- to IgG-cell ratio and another with a ratio of less than 1. They proposed that in some patients with pernicious anaemia, a strikingly reduced proportion of IgA immunocytes could represent a primary defect predisposing to mucosal damage. They showed in fact that the reduced ratio could be ascribed to a decrease of the IgA- and an increase of IgG-cell density in the lamina propria. It should be noted, however, that a decreased cell density might be caused by oedema; a better reflection of the local immunological activity is obtained by cell enumeration based on a mucosal tissue unit as carried out in the present study. Nevertheless, regardless of the quantitative method used, a reduced IgA- to IgG-isotype ratio reflects an altered immunological homeostasis in the mucosa.

The biological significance of locally produced gastric IgM is difficult to evaluate. Because IgM can be transported by SC-producing epithelium, it may contribute to external defence although being less stable than secretory IgA. The number of IgM-producing cells varies considerably among different secretory tissues, the highest proportion being found in the proximal small intestine. It is unknown whether such variations in local isotype profiles may be ascribed to dissimilar precursor pools of B cells.

### Table 2 Immunohistochemical studies of Ig-producing cells in human gastric mucosa

<table>
<thead>
<tr>
<th>Gastric region</th>
<th>Histological classification</th>
<th>Cell quantitation</th>
<th>Percentage distribution</th>
<th>IgA/IgG ratio</th>
<th>Reference</th>
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<td></td>
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<td>IgA</td>
<td>IgM</td>
<td>IgG</td>
<td>IgA</td>
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<td>Normal and low-grade gastritis</td>
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<td>24-2</td>
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*Number of cells per mm² of interstitial area. †Patients with pernicious anaemia. ‡Number of cells per mm² of lamina propria. §Number of cells per area of lamina propria. ||IgE cells=3%. ¶Exclusively IgD and IgE cells. §§IgE=12%. ‖IgE=7-5%.
Immunocytes in gastric mucosa

homing to the different sites or reflect region-related topical antigenic or mitogenic effects on local B-cell differentiation. We found a relatively high proportion of IgM cells in the gastric mucosa, although it was much lower than that reported by Camilleri et al. As IgM has complement-activating properties, its enhanced local production in gastritis may well be of immunopathological importance.

A striking preponderance of IgA immunocytes was found in gastric mucosa, signifying an active secretory immune system. With increasing grade of gastritis, the total number of immunocytes was significantly increased but disproportionately so for the IgG isotype. Enhanced local production of IgG probably reflects a second line of defence which over time, however, through immunopathological mechanisms may contribute to glandular atrophy and perpetuation of gastritis. The subclasses of IgG included in the gastric immune responses are unknown and will be mapped in further immuno- histochemical studies by means of monoclonal antibodies.

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