Studies of intestinal lymphoid tissue. XIV-HLA status, mucosal morphology, permeability and epithelial lymphocyte populations in first degree relatives of patients with coeliac disease


Abstract
Fifty two first degree relatives of patients with coeliac disease were investigated for HLA status, small intestinal permeability, and mucosal morphology together with the size of the intraepithelial lymphocyte pool and indices of lymphocyte activation, in an attempt to identify genetically determined markers of the disease. Thirty eight per cent of these subjects had increased intraepithelial lymphocyte populations and a highly significant association with HLA-DR3 compared with controls. Their intestinal permeability to chromium-labelled ethylenediamine tetaacetate was invariably normal and there was no evidence of abnormal mucosal architecture, increased crypt cell mitotic activity or lymphocyte 'activation'. Although increased intraepithelial lymphocyte counts clearly do not cause alterations in intestinal structure or function, it is likely that this parameter together with the HLA-DR3 status identifies a genetically determined predisposition to the disease which may only become clinically evident with larger doses of ingested gluten.

Despite considerable investigation over many years, the pathogenesis of coeliac disease remains unexplained. One of the major obstacles to progress is that after a gluten free diet, most detectable abnormalities revert to normal, while in a few instances in which some kind of dysfunction remains, it is extremely difficult to exclude continued inadvertent ingestion of gluten, or perhaps, the effect of irreversible damage caused by long standing mucosal inflammation.

In order to overcome some of these problems and to permit study of the early initiating pathogenic factors in this condition, one approach is to study well treated patients exposed to gluten for relatively short times. An alternative is to study apparently unaffected close relatives of known propositi in order to identify isolated inherited abnormalities that do not by themselves lead to full expression of the disease, perhaps because of lack of other necessary interacting genetic, or environmental, factors.

Based on the latter approach, we have carried out a detailed investigation of a large number of first degree coeliac relatives who had been closely followed up in North West England for 10-20 years. The aim of this collaborative study was to compare the HLA-DR status of such individuals with small intestinal permeability, mucosal morphology, the size of the intraepithelial lymphocyte pool and indices of epithelial lymphocyte 'activation', of which some may have importance in the pathogenesis of coeliac disease.

Methods
First Degree Relatives of Patients with Coeliac Disease
Of approximately 200 relatives from 28 families containing well documented coeliac disease propositi, 52 first degree subjects were selected, based on jejunal biopsies suitable for morphometric quantitation by computerised image analysis. They comprised 23 men (mean age 37 years; range: 18-78) and 29 women (mean age 42 years; range: 17-72). Those relatives with clinically symptomatic, or inapparent (latent), coeliac disease were, however, excluded from further study.

The majority of the 52 first degree relatives were healthy subjects without any significant previous medical history. Five had complained of recurrent mouth ulcers, another was on thyroxine replacement therapy and two had non-insulin dependent diabetes mellitus.

Apart from one individual with a previous history of diarrhoea, at the time of study, none had any gastrointestinal disturbance. For the 23 men and 29 women, respectively, there was no significant difference (Mean (SD)) in concentrations of haemoglobin, 14-4 (1-2) and 13-0 (1-8) g/l; serum folate, 6-4 (2-5) and 7-2 (4-8) µg/l; serum iron, 14-8 (3-1) and 16-6 (9-9) µmol/l; serum calcium, 2-4 (0-16) and 2-4 (0-13), or serum IgA, 25 (9) and 20 (6) g/l.

Control Subjects
(a) For evaluation of morphometric data, results were compared with those obtained for (i) 14 untreated coeliac patients and (ii) a group of 10 patient-control subjects referred for diagnostic gastrointestinal investigation to Department of Medicine at Hope Hospital: none of the latter was considered to have coeliac disease and jejunal morphology was judged normal by accepted criteria. (b) HLA-DR locus frequencies in the first degree relatives were compared with a sample of blood donors and members of staff from the same geographic area, as detailed below.
HISTOLOGICAL TECHNIQUE

Jejunal biopsies were taken with a Watson capsule from the duodenal-jejunal flexure, quickly orientated, spread out on card, flooded with buffered formalin (pH 7-2), embedded in paraffin wax, sectioned at 5 μm thickness in a plane perpendicular to the mucosal surface and stained with H&E.

Eight to 10 serial sections were mounted per slide: alternate sections, provided their plane of sectioning was well oriented, were used for morphometric analysis. For this purpose, chosen sections were projected through a high resolution colour television (Sony) camera to a MOP-Videoplan® (Kontron-Reichert, Slough, England) computerised image analysis system. Photomicrographs were taken on Ilford (Ilford Ltd, Ilford, Essex, UK) 35 mm film as desired, and enlarged appropriately.

QUANTITATIVE MORPHOMETRY

1: Surface and crypt epithelial volume

In order to determine surface and crypt epithelial volumes (V_SE; V_CR) per specimen, profiles of relevant sectioned epithelium were outlined by the scribing cursor together with the length of underlying muscularis mucosae. For valid comparison, measurements were standardised to a constant test area (100×100=10^4 μm^2) of muscularis mucosa. The relative coefficients of variation for each series of measurements per specimen were <10%; these calculations are automatically included in the print-out of the measuring programme supplied by the manufacturers.

Total crypt cell mitotic figures per specimen, as a reflection of crypt cell proliferative activity, were also quantitated with reference to 10^4 μm^2 muscularis mucosa.

2: Size distribution of intra-epithelial lymphocytes

Based on previous work, the mean size of intraepithelial lymphocytes per specimen was expressed in terms of nuclear diameters. The circumference of nuclear profiles was traced with the cursor from which crude diameters were computed. Sufficient observations to ensure a constant mean (SD) were incorporated: on average, 80–100 lymphocyte nuclear profiles per specimen were required to meet these criteria.

The distribution of the crude profile diameters was corrected for 'lost' profiles, and the mean of this revised distribution was further corrected for imperfect 'non-sagittal' sectioning by multiplying by 4/π. In this way, the true nuclear diameter (D_N) of a representative sample of intraepithelial lymphocytes per specimen was obtained.

3: Determination of intraepithelial lymphocyte population size

In order to calculate the actual number of cells (N) within a defined tissue volume (V), the mean diameter (such as D_N) must be calculated in order to determine 'effective section thickness' (EST) which is determined from the relationship (t + D_N) where t is actual section thickness (5 μm). The absolute number of intraepithelial lymphocytes contained within an epithelial volume (V_SE or V_CR) with respect to a 10^4 μm^2 test area of muscularis mucosa was determined by enumerating lymphocyte profiles within surface, or crypt, epithelium overlying a length of muscularis equivalent to [10 000/D_N + 5]μm. Nuclear profiles were preferred in calculating EST because they are more usually circular in outline compared with cytoplasmic profiles, and thus more accurately measured.

The proportion of 'immunoblastoid' intraepithelial lymphocytes with D_N > 6 μm was per specimen was calculated from the distributions obtained in (2) above. The number of metaphase-figures counted in a total of 3000 epithelial lymphocytes per specimen was expressed as per cent mitotic index.

INTESTINAL PERMEABILITY

Of the 52 subjects included in this study, 26 gave informed consent to undergo an intestinal permeability test with 51Cr-EDTA, assessed by determining the 24 h urinary excretion of the probe molecule as described previously. Ethical approval for the use of this test was given by the appropriate local research committees.

HLA TYPING

HLA-DR typing was performed by a cytotoxicity technique based on an adaptation of a two-colour fluorescence method. The panel of available sera defined seven specificities (DR1-DR7). The controls were 168 staff members or blood donors from the same geographic areas.

STATISTICAL ANALYSIS

Statistical analyses were performed with the SPSS package. Group means were compared with the Mann-Whitney U test, a p value of <0·05 being taken as significant. The Spearman correlation coefficient was used to compare epithelial lymphocyte numbers with (i) surface epithelial volume or (ii) 51Cr-EDTA urinary excretion. Differences between antigen frequencies were examined by Fisher's exact (two-tailed) test as previously described.

Results

INTRAEPITHELIAL LYMPHOCYTE POPULATIONS (N_{V,SE}; N_{V,CR}):

Values for N_{V,SE} were positively skewed for relatives of patients with coeliac disease (skewness +1.18, which differed significantly (p < 0·01) from zero). Thus all values were logarithmically transformed, the normalised distribution having a recalcualted skewness of −0·10 (p = NS).

In comparison with N_{V,SE} for 20 disease controls, the 52 first degree coeliac disease relatives were arbitrarily divided into subgroups 1 and 2. In group 1, N_{V,SE} fell below the upper
95% confidence limits of the control sample (240 ± 93) while in group 2, comprising 38% relatives studied, \( N_{v, SE} \) exceeded these limits (630 (122)) as shown in representative micrographs (Fig 1). Corresponding values for absolute intraepithelial populations in the crypts (\( N_{v, CR} \)) were, for group 1 subjects 21 (8), and for group 2 subjects with raised villous epithelial lymphocyte populations, 34 (18) (differences NS). Neither was there any difference in crypt cell mitotic activity (per \( 10^4 \) \( \mu m^2 \) muscularis mucosae) between group 1 (6-3 (3-3)) and group 2 (6-6 (4-5)) individuals. Thus the increment in intraepithelial lymphocytes was confined to villus epithelium only (group 2 relatives).

**SIZE DISTRIBUTION AND MITOTIC ACTIVITY OF INTRAEPITHELIAL LYMPHOCYTES**

There was no difference in the mean nuclear diameter (\( D_{N} \)) of intraepithelial lymphocytes between the disease-controls (5-0 (0-2) \( \mu m \)) and coeliac disease relatives (4-6 (0-3) \( \mu m \)). Mean values for \( D_{N} \) between group 1 (4-5 (0-3) \( \mu m \)) and group 2 (4-4 (0-3) \( \mu m \)) coeliac disease relatives were also identical (Fig 2). The percentage of large lymphocytes (\( D_{N} > 6 \mu m \)) in the disease-controls and in each subgroup of coeliac disease relatives were similar and considerably less than found in patients with untreated coeliac disease**"** (Fig 2). The mitotic activity (sample of 3000 intraepithelial lymphocytes per specimen) determined for each subgroup of coeliac relatives did not exceed the 0-2% seen in patients with untreated coeliac disease.**"** Thus, based on size and mitotic activity, there was no evidence of lymphocyte 'activation' within either group of coeliac relatives.

**Figure 1: Right hand panel shows a marked increase in small, non-mitotic intraepithelial lymphocytes (EL) within normal villi (arrows in a group 2 CD relative), compared with appearances in a group 1 mucosa (left hand panel). (Magnifications: × 1925: Insets, ×67).**

**Figure 2: Mean nuclear diameters of intraepithelial lymphocytes (\( D_{n} \), Lyt – left hand axis) and the percentage \( D_{N} > 6\mu m \) (right hand axis) in controls (C), CD relatives (R) and their subgroups, 1 and 2, compared with a group of untreated coeliacs (CD). The coeliac relatives either considered together, or as subgroups based on lymphocyte populations, do not differ from the controls, whereas untreated coeliac patients display 'activated' EL with an increased mean nuclear diameter, and increased proportion >6\( \mu m \) diameter.**
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The percentage 24 h urinary excretion of \(^{51}\text{Cr}-\text{EDTA}\) in the 26 relatives who were tested showed a wide range of values, from 0.8 to 4.7.

**Figure 4:** For the entire group of relatives (left hand panel) studied there is no significant correlation between absolute number of epithelial lymphocytes \((NV_{SE})\) and surface epithelial volumes \((V_{SE})\). There is no significant relationship between epithelial lymphocyte populations \((NV_{SE})\) (right hand panel) and percentage excretion of \(^{51}\text{Cr}-\text{EDTA}\) in the 26 relatives tested.
exclude all such individuals from this study. In comparison with previous studies, however, we were left with a significant proportion (38%) of first degree relatives, predominantly of DR3 phenotype, whose mucosa revealed a markedly increased intraepithelial lymphocyte population. Furthermore, these individuals were identified in every pedigree examined, and not restricted to one-third of all families as is characteristic for fully developed ‘flat’ mucosal lesions, whether associated with symptoms or not. If all our group 2 first degree relatives were to progress and develop overt disease, then the familial prevalence rate for coeliac disease in the North-West region would considerably exceed the consistent ± 10% rate recorded in the UK, 8,10 Irish Republic, 9 USA11,12 and other parts of the world. 8,10 It is therefore evident that these individuals appear to have a non-progressive mucosal abnormality.

The significance of the lymphoid infiltrate in group 2 relatives is unclear. They were assumed to be ingesting a normal diet unlike experimental subjects 13,14 in whom gluten loading may result in an accumulation of intraepithelial lymphocytes. What is evident from this study is that the presence of an increased number of intra-epithelial lymphocytes is neither causally related to increased mucosal permeability, nor to any structural abnormality. In this respect these observations parallel earlier studies when we fed treated coeliac patients with small, graded doses of Frager’s fraction III, resulting in dose dependent rises in intraepithelial lymphocytes, with or without accompanying mucosal deterioration: similarly, these lymphocytes were small and non-mitotic. 15 This is in keeping with other data suggesting that gluten induced accumulation of lymphocytes within epithelium plays no role in mucosal damage, because (i) intraepithelial lymphocytes may increase in number after dietary gluten restriction 16 * (ii) crypt cells in untreated coeliac disease, despite a five-fold increase in intra-epithelial lymphocytes, 17 are evidently unimpaired in their hypertrophic response or ability to generate goblet, argentaffin and Paneth cells and (iii) many gluten sensitive individuals with dermatitis herpetiformis carry raised lymphoid infiltrates in their villi without evidence of functional or structural enteropathy. Collectively, all these observations suggest that intraepithelial lymphocytes alone are unlikely to play a role in initiating, or sustaining, mucosal damage throughout the entire ‘gluten-sensitivity spectrum’. It would be interesting to challenge such relatives with gluten, however, to confirm that they do indeed have a subclinical form of the disease, and that mucosal flattening would necessarily ensue.

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