FAS engagement drives apoptosis of enterocytes of coeliac patients

L. Maiuri, C. Ciacci, V. Raia, L. Vacca, I. Ricciardelli, F. Raimondi, S. Auricchio, S. Quarantino, M. Londei

Abstract

Background—Villus atrophy is the most distinctive sign of untreated coeliac disease (CD) and epithelial apoptosis is considered to be involved in this stage of the coeliac lesion. The extent of villus atrophy is, however, not homogeneous and patients with patchy or mild lesions have been described.

Aims—To address: (a) the degree of “patchiness” in untreated CD patients; and (b) to clarify if apoptosis, and eventually which trigger drives it, causes epithelial damage.

Patients—Twenty of 40 untreated, 14 treated coeliac patients, and 15 controls received five or more multiple duodenal biopsies; the remaining 20 untreated CD patients had no more than three biopsies.

Methods—All biopsies were analysed to monitor the presence of a “flat” mucosa. Biopsies of 14 untreated, 10 treated coeliacs, and seven controls were cultured with or without gliadin. DNA fragmentation was studied by terminal deoxynucleotidyl transferase (TdT) mediated dUTP digoxigenin nick end labelling (TUNEL), and FAS and Ki67 expression by immunohistochemistry. Antiendomysium antibodies (EMA) were surveyed in biopsy culture supernatants.

Results—A pattern of patchy duodenal lesions was observed in all untreated CD patients biopsied up to five times. High enterocyte FAS expression, and a high number of TUNEL+ and Ki67+ enterocytes were detected in areas with villus atrophy but not in those with a normal morphology (p<0.001). Conversely, EMA in culture supernatants and signs of immunological activation were present in all untreated CD biopsies. In vitro gliadin challenge increased the number of TUNEL+ and Ki67+ enterocytes (p<0.001 v cultures with medium alone) in “flat” biopsies. Neutralising anti-FAS monoclonal antibodies were found to control gliadin induced enterocyte apoptosis (p>0.01) while agonist anti-FAS monoclonal antibody increased it (p<0.001).

Conclusions—Patchy lesions are observed in untreated CD mucosa and epithelial FAS engagement is a key trigger in driving villus atrophy in CD.

Keywords: apoptosis; FAS; enterocytes; coeliac disease

Coeliac disease (CD) is a frequent and, to date, under diagnosed disease of the small intestine characterised by the presence of endomysium antibodies (EMA) in the sera of untreated CD patients. Three of the major histological features of the small intestinal mucosa of untreated CD patients are: gliadin dependent activation of supposed pathogenic T cells; T lymphocyte intraepithelial infiltration; and crypt hyperplasia with villus atrophy. The latter, the “flat” mucosa, is possibly the most canonical feature of CD although it cannot be considered in isolation but in the wider context of a remodelling process of the small intestine. It is now accepted that CD is not limited to patients with severe enteropathy but can also be applied to patients with minor pathological changes, where “patchy” lesions are sometimes observed. Evidence of patchy duodenal lesions in the same CD patient provided an intriguing descriptive study of mucosal damage in CD but an explanation for these unexpected findings is still awaited. Here, we analysed a larger sample of CD patients by multiple endoscopic biopsies to fully evaluate the extent of the patchy lesion profile and also to unravel the mysteries controlling induction of the flat mucosa.

It has previously been reported that DNA fragmentation is observed in the majority of enterocytes of untreated CD intestine and that after gluten withdrawal, a normal profile is restored. One of the most studied pathways leading to apoptosis is by FAS engagement, the prototype of the “death” receptors widely expressed by many cell types which we have recently reported to be rapidly upregulated in treated coeliac duodenum on in vitro challenge with gliadin.

Analysis of patchy lesions in coeliac duodenum represents an ideal tool to clarify the correlation among FAS expression, enterocyte apoptosis, crypt epithelial proliferation, and villus atrophy. Moreover, using the organ culture model of coeliac duodenal explants, we tested the true role of FAS engagement in driving epithelial apoptosis and villus flattening in CD using agonist or antagonist anti-FAS monoclonal antibodies (mAbs).

Our study indicates that a definite correlation between villus flattening, enterocyte FAS expression, apoptosis, and crypt cell proliferative-

Abbreviations used in this paper: CD, coeliac disease; EMA, endomysium antibodies; mAb, monoclonal antibody; Ig, immunoglobulin; PT, peptic-tryptic; TUNEL, terminal deoxynucleotidyl transferase (TdT) mediated dUTP digoxigenin nick end labelling; IEL, intraepithelial lymphocytes.
tion is present in coeliac duodenum. Moreover, our data clearly suggest that FAS engagement plays a key role in induction of epithelial damage and that the difference between patchy lesions and a normal histology is the specific lack of expression of FAS. Hence our studies may help in the understanding of the precise correlation that exists between induction of damage and the remodelling process observed in CD.

Methods

PATIENTS

Forty untreated CD patients (mean age 31.2, range 3–54) underwent duodenal endoscopy and biopsy for diagnostic purposes. In all patients serum anti-EMA were detected. Twenty patients underwent less than three biopsies which all revealed villus atrophy on routine histological examination. The other 20 patients underwent at least five biopsies, examined immediately at the dissecting microscope. In 12 of these 20 patients the first sample analysed showed villus flattening whereas in eight patients the first sample showed apparent normal villi. Analysis at the dissecting microscope of all fragments obtained by multiple endoscopic biopsies revealed a patchy duodenal lesion with various extension of the villus atrophy along the duodenum in all 20 patients. For each patient, one fragment with villus flattening and one sample with apparent normal villi were embedded in OCT and used for histological analysis to confirm the appearance observed at the dissecting microscope. The other fragments (with villus atrophy or normal villi) were divided into two parts each and cultured as described below.

Fourteen of the untreated patients (mean age (range) 37.3 (21–53) years) were studied after a strict gluten free diet spanning a period of at least 12 months when they were rebiopsied (termed treated CD patients). All were EMA negative at the time of the second biopsy.

Fifteen non-CD patients (mean age (range) 31.1 (14–60) years), undergoing duodenal endoscopy because of oesophagitis (n=5), gastritis (n=5), or chronic non-specific diarrhoea (n=5) had multiple biopsies. In all biopsies, all fragments showed normal villus architecture at the dissecting microscope as well as after histological analysis. Informed consent was obtained from all CD patients and controls.

IN VITRO ORGAN CULTURE STUDIES

Duodenal explants from duodenal regions with villus atrophy from 10 untreated CD patients were cultured in vitro9–11 in the presence or absence of a PT gliadin digest (1 mg/ml) or agonist anti-FAS CH-11 mAb. Duodenal explants from 10 treated CD patients were cultured in vitro for 24 hours9–11 in the presence of culture medium with or without a PT digest of gliadin (1 mg/ml) (in all cases), and with maize prolamin peptides (1 mg/ml) (in six cases) or agonist anti-FAS CH-11 mAb (in four cases). In seven controls, in vitro cultures with or without PT gliadin digest (1 mg/ml) were performed; in three cases agonist anti-FAS CH-11 mAb was also tested.

DETECTION OF DNA FRAGMENTATION

DNA fragmentation on tissue sections was assayed in an experimental group previously reported8 by terminal deoxynucleotidyl transferase (TdT) mediated dUTP digoxigenin nick end labelling (TUNEL). In CD subjects with patchy lesions, DNA fragmentation was assessed in samples with villus atrophy as well as in those with a normal mucosal architecture. Specificity control experiments were performed as previously reported.10 Two colour immunofluorescence was used to determine whether apoptotic cells were T cells or epithelial cells. The experiments were carried out by developing the TUNEL reaction with phycoerythrin (PE) conjugated anti-digoxigenin mAb (Boehringer, Mannheim, Germany) and simultaneously incubating tissue sections with anti-CD3 mAb (1:100; Dako, Copenhagen, Denmark) followed by FITC conjugated antimouse Ig (1:30; Dako).

IMMUNOHISTOCHEMISTRY

Antigen detection on frozen tissue sections was performed by immunohistochemistry as previously reported8 with mAbs anti-Ki67 (1:25; Dako) or CD3 (1:100; Dako), anti-FAS (M3 and M38) (mouse IgG1, 1:30; Immunex Corp.), CD25 (1:30; Dako), and ICAM1 (1:400; 509 Ylem CVA) by peroxidase or alkaline phosphatase staining techniques according to a method previously described. Anti-FAS-ligand mAbs (A11 and H11 rat biotinylated Ig1:60; Alexis-Corporation, San Diego, California, USA) were used to detect FAS-ligand expression in biopsies from CD patients. The antibodies were visualised using a peroxidase staining technique following by a method previously described.1 Anti-FAS ligand mAbs (A11 and H11 rat biotinylated Ig1:60; Alexis-Corporation, San Diego, California, USA) were also used to detect FAS-ligand expression in biopsies from CD patients. The antibodies were visualised using a peroxidase staining technique following by a method previously described. Numbers of intraepithelial lymphocytes (IEL) were counted as CD3+ cells/100 enterocytes8 and Ki67+ enterocytes as percentage of crypt enterocytes. Staining of epithelial cells by anti-FAS mAb was arbitrarily graded from 0 to +2, based on the intensity of experimental conditions. Neutralising anti-FAS M38 mAb was also tested in three cultures and the results correlated with those obtained using neutralising anti-FAS M3 mAb. Control antibodies anti-CD80 (5 µg/ml) (in four cases) or anti-human lactase mAb mlac1 (5 µg/ml)15 (in four cases) as well as CTLA4-immunoglobulin (Ig) (Genetic Institute Corporation, Cambridge, USA) (in five cases) were also added to gliadin.
staining in more than 70% of cells (undetectable=0, low=1+, intense=2+). Guidelines for the scoring system were established at the start of the study and four observers independently analysed the samples; the results were compared afterwards.

At least five slides for each sample were blindly evaluated. Specificity control experiments were performed using mouse IgG or IgM against inappropriate blood group antigens, and simultaneously analysing different cultured samples belonging to the same individual.

DETECTION OF EMA IN CULTURE SUPERNATANTS

EMA were detected in culture supernatants of cultured untreated CD biopsies (10 from areas with villus atrophy and four from areas with normal histology) after incubation with medium alone, in accordance with methods described previously.11

STATISTICAL ANALYSIS

Samples from each category were compared. The Student’s t test for paired samples was used to discriminate between untreated CD intestine TUNEL+ enterocytes and Ki67+ enterocytes after challenge with medium alone, with medium supplemented with gliadin, or with gliadin supplemented with mAbs. Non-parametric tests (Wilcoxon test) were also applied and the results compared with those obtained using parametric tests. Fisher’s test was used to compare tissues with undetectable or low FAS expression (0 to 1+) with those showing intense (2+) expression. Fisher’s test was used to compare FAS expression in tissue specimens after gliadin challenge with those after challenge with medium alone.

Results

PATCHY MUCOSAL LESIONS ARE FREQUENTLY OBSERVED IN UNTREATED CD DUODENUM

In all samples, duodenal mucosal findings obtained using the dissecting microscope were confirmed by histological analysis. In the 20 untreated CD biopsies in which more than five multiple biopsies were performed, we observed

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Table 1  Overall representation of the characteristics of the mucosal manifestations in coeliac disease (CD) and controls

<table>
<thead>
<tr>
<th></th>
<th>Untreated CD</th>
<th>Normal mucosa</th>
<th>Treated CD</th>
<th>Controls</th>
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</thead>
<tbody>
<tr>
<td>Villus atrophy</td>
<td>High</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>Enterocyte apoptosis</td>
<td>High</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
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<tr>
<td>Enterocyte FAS expression</td>
<td>High</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>IEL FAS-ligand expression</td>
<td>High</td>
<td>High</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>LPMNC activation</td>
<td>High</td>
<td>High</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>EMA in culture supernatant</td>
<td>Present</td>
<td>Present</td>
<td>Absent</td>
<td>Absent</td>
</tr>
</tbody>
</table>

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Figure 1  DNA fragmentation and Ki67 expression in coeliac disease (CD) and control intestine. (A) DNA fragmentation in enterocytes of CD and control intestine. (B) Expression of Ki67 in crypt enterocytes of CD and control intestine (mean (SD)). Untreated CD, untreated biopsies with patchy mucosal lesions (n=20); treated CD, treated biopsies (n=14); controls, biopsies from non-coeliac patients with a normal villus architecture (n=15). ***p<0.0001 v treated CD and controls. (C) Pattern of intraepithelial DNA fragmentation in untreated CD duodenum. DNA fragmentation (TUNEL technique, red colour) is absent in the nucleus of all intraepithelial lymphocytes (CD3, green colour), while it is detected in the majority of enterocytes. Original magnification ×240. Two colour immunofluorescence: TUNEL technique, RPE labelling (red); immunohistochemistry, CD3 labelling (green) (see methods).
that the flat area corresponded in all cases to villus atrophy and crypt hyperplasia, and in 18/20 samples increased IEL counts (above 40/100 enterocytes) were detected. Well represented vili observed with the dissecting microscope corresponded to normal villus length with a villus height/crypt depth ratio higher than 3,7 and IEL counts greater than 40/100 enterocytes were found in only 13/20 samples. In all untreated CD samples, regardless of mucosal damage, common features of intestinal immune activation9−10 were found in the lamina propria (number of CD25+ and ICAM1+ cells in the lamina propria higher than mean +3 SD of control values), as previously reported by us7 (table 1).

DNA FRAGMENTATION, AND KI67 AND FAS EXPRESSION IN ENTEROCYTES CORRELATE WITH MUCOSAL LESIONS

Untreated CD samples with villus atrophy

In untreated CD biopsy samples with villus atrophy, TUNEL positivity was increased compared with controls (p<0.0001) (fig 1A) and expression of Ki67 was strongly augmented in crypts, in agreement with previous reports7 (p<0.0001 v controls) (fig 1B). Virtually no CD3+ cells stained TUNEL positive, indicating that all apoptotic cells were epithelial. Interestingly, some TUNEL positive (red) epithelial cells were not near to CD3+ (green) (fig 1C). This finding may indicate that not all epithelial death is caused by infiltrating FAS-ligand positive IEL and that other mechanisms may be involved, although no definitive conclusions can be drawn at present. Intense FAS expression (2+) in surface and crypt enterocytes was found in 16/20 samples. In these samples expression of FAS was detected in the majority (more than 70%) of enterocytes (v 0/15 in controls; p<0.01) (table 1).

FAS-ligand expression was detected in IEL (more than 50%) of all but four untreated CD biopsies with villus atrophy (data not shown). Interestingly, the pattern of FAS-ligand expression was highly reminiscent of that described by Viard et al in patients with Lyell’s syndrome.16

Untreated CD samples with normal morphology

In all untreated CD samples with a normal mucosal architecture, the pattern of DNA fragmentation (fig 1A) and Ki67 antigen expression (fig 1B) was similar to controls (p>0.5). Also, high FAS expression was not found in any of the eight cases (p>0.5 v controls). In a few cases we detected a sharp change in expression of FAS at the border between flat (high) and normal (low) areas (fig 2).

FAS-ligand expression in untreated CD intestine with normal villus architecture was similar to that found in untreated CD with villus atrophy (table 1).

Treated CD samples

In all treated CD samples the pattern of DNA fragmentation (fig 1A) and Ki67 antigen expression (fig 1B) was similar to that observed in controls (p>0.5). High enterocyte FAS expression was observed only in 1/14 cases (p>0.5 v controls). In these samples no evidence of immunological activation was observed (table 1), as previously reported.7 IEL in treated CD and controls were virtually all FAS-ligand negative (table 1).

ORGAN CULTURE STUDIES

In untreated CD biopsies with villus atrophy, gliadin induces an increase in enterocyte apoptosis via a FAS dependent pathway. Increased expression of FAS in enterocytes did not automatically mean that FAS engagement was crucial in driving epithelial apoptosis. We used in vitro culture studies to verify whether or not expressed FAS could have delivered a functional signal.

In seven cultured untreated CD samples with villus atrophy in which FAS expression was intense before in vitro manipulation, 24 hour gliadin challenge caused an increase in the number of TUNEL positive enterocytes compared with the profile observed with medium alone (p<0.001) (fig 3A). Expression of Ki67 was markedly decreased after incubation with medium alone (p<0.0001) whereas gliadin induced Ki67 expression observed before in vitro manipulation (p>0.5) (fig 3B).

Both neutralising anti-FAS antibodies were effective in reducing the number of TUNEL+ enterocytes compared with the profile observed in vitro after incubation with gliadin alone (p<0.001) (fig 3A, C, D) while they failed to control Ki67 expression (data not shown). No protective effect was observed after incubation with anti-human lactase mIac117 or anti-CD80 mAbs, which bear the same isotype of the neutralising anti-FAS antibody in four tested cases (p>0.5) (fig 3A). In five of the cultured untreated CD biopsies with villus atrophy, CTLA4-Ig, a powerful and specific inhibitor of T cell activation,19 was unable to control the gliadin induced increase in DNA fragmentation of enterocytes (p>0.5) (fig 3A). However, CTLA4-Ig downregulated the pattern of mucosal immune activation (data not shown) indi-
cating its efficacy in controlling the immuno-
mediated gliadin induced activation. In four
cultured untreated CD samples with villus atro-
phy, agonist CH-11 anti-FAS mAbs induced a
dramatic increase in TUNEL+ enterocytes and
derangement of the mucosal architecture
(p<0.001) (fig 3A, E) while it did not increase
Ki67 expression (data not shown) (table 2).

In untreated CD biopsies with normal villus
architecture, low enterocyte FAS expression was
able to launch the apoptotic programme.

In the four cultured untreated CD samples with
normal villus architecture in which FAS
expression was low to absent before in vitro
culture, triggering of the FAS receptor with
agonist CH-11 anti-FAS mAbs was not
effective in inducing enterocyte apoptosis
(p>0.5 v cultures with medium alone) (fig 4A).
A 24 hour gliadin challenge was similarly
unable to increase the number of TUNEL+
enterocytes (fig 4A) or to increase Ki67
expression in crypts (fig 4B) (p>0.5 v cultures
with medium alone). Moreover, in these biop-
sies, no increased expression of enterocyte FAS
receptor was induced by gliadin challenge in
any of the four cultured biopsies (p>0.5 v cul-
tures with medium alone) (table 2).

Untreated CD biopsies release EMA in culture
supernatants regardless of duodenal morphology
In all cultured untreated CD samples with vil-
lus atrophy, analysis of culture supernatants'1
revealed spontaneous release of EMA in the

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**Figure 3** In vitro organ culture of untreated coeliac disease (CD) intestine with villus atrophy: DNA fragmentation and
Ki67 expression after 24 hour challenge with medium or gliadin. Effect of neutralising anti-FAS M3 or agonist anti-FAS
CH-11 monoclonal antibodies (mAbs) on enterocyte apoptosis. (A) DNA fragmentation and (B) Ki67 expression before
and after in vitro culture. Untreated CD samples with villus atrophy before in vitro culture (n=7) and after 24 hour
challenge with medium alone (n=7), gliadin (n=7), gliadin supplemented with anti-FAS M3 mAb (n=7), or medium
supplemented with anti-FAS CH-11 mAb (n=4). The number of TUNEL+ enterocytes (mean (SD)) are given as
percentage of both surface and crypt enterocytes and Ki67+ enterocytes (mean (SD)) as percentage of crypt enterocytes.

**p<0.001 v samples after 24 hour challenge with medium alone; †††p=0.001 v samples after 24 hour gliadin challenge.
(C) Pattern of DNA fragmentation after 24 hour gliadin challenge. Note that the majority of enterocytes are TUNEL+;
staining is also evident in many lamina propria mononuclear cells. (D) Pattern of DNA fragmentation after 24 hours of in
vitro challenge with gliadin supplemented with neutralising anti-FAS M3 mAb. The number of TUNEL+ enterocytes is
decreased compared with the pattern observed after gliadin challenge. (E) Pattern of DNA fragmentation after 24 hours of
in vitro challenge with medium supplemented with agonist CH-11 anti-FAS mAb. A marked increase in the number of
TUNEL+ enterocytes with derangement of epithelial architecture is evident. Original magnification: C, D and E (×180);
TUNEL technique, peroxidase staining.
absence of gliadin stimulation, in accordance with previous reports. Interestingly, spontaneous release of EMA was detected in culture supernatants even in all four cultured untreated biopsies with normal mucosal architecture (table 2).

In treated CD samples, gliadin induces FAS expression in enterocytes

In all 10 cultured treated CD samples, 24 hour gliadin challenge was unable to increase the number of TUNEL+ or Ki67+ enterocytes (data not shown) while it was effective in enhancing enterocyte FAS expression in 7/10 cases (p<0.01), as described in a previous report (fig 5A–5C) (table 2).

Discussion

Complex patterns of events control the evolution of the mucosal manifestations of CD, which is characterised by clear signs of mucosal damage and concomitant features of tissue remodelling. The final result of this destructive/remodelling activity is believed to be an evenly distributed “flat” mucosa with an overall increase in the whole mucosa with crypt enlargement. This pathological pattern is customarily believed to be caused by immunological recognition of gliadin. However, it is recognised that patchy lesions are observed in some coeliac patients but the proportion with a patchy pattern is not clearly defined and is considered to be infrequent. This study provides several significant findings and clarifies some of the pathogenic features of CD. In the first instance, we observed that patchy lesions are frequently observed in untreated CD mucosa and that they are not the exception in untreated CD duodenum. This may have an important clinical significance as diagnosis of CD is still widely based on small intestine histology. We found that patchiness was not rare and conversely quite frequent in untreated CD, and hence some patients could be wrongly diagnosed as not having CD.

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Overall representation of the characteristics of the mucosal manifestations in organ culture of the small intestine of coeliac disease (CD) patients and controls</th>
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<tbody>
<tr>
<td>Vector</td>
<td>Untreated CD</td>
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<tr>
<td>Increase in enterocyte apoptosis*</td>
<td>Yes</td>
</tr>
<tr>
<td>Increase in enterocyte Ki67 expression*</td>
<td>Yes</td>
</tr>
<tr>
<td>Increase in enterocyte FAS expression*</td>
<td>Yes</td>
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</table>

*Organ cultures with gliadin digest cultures with medium alone.
With regard to the pathogenic factors controlling induction of the epithelial phase, the “flat” patches presented with clear signs of epithelial apoptosis and proliferation, as observed by others, high expression of FAS, evidence of in situ immunological activation, and spontaneous release of EMA. Analysis of patches with a “normal” structure indicated that there were no signs of epithelial involvement (proliferation, apoptosis, or FAS expression) while immunological activation and EMA production were detected in untreated CD patients. Thus we have an apparent lack of connection between immunological activation and mucosal damage. Thus these results suggest that the epithelial phase of CD has an apparently more complex mechanism of induction and that it might not be driven solely by immunological recognition of gliadin. Expression of FAS seems to characterise the “flat” regions of the patches and more importantly, FAS engagement has a key role in induction of mucosal damage driven by gliadin challenge. The study using neutralising or agonist anti-FAS antibodies unequivocally indicated that gliadin induced FAS modulation and its engagement has a pivotal role in epithelial damage. Even in this situation it appears that there is a lack of connection between immunorecognition of gliadin and epithelial damage, as we indicated under other experimental conditions. Indeed, CTLA4-Ig, an extremely efficient device to control T cell activation, could not control induction of apoptosis by gliadin in untreated CD explants.

An intriguing finding was that in untreated CD, duodenal regions of “normal” mucosa were contiguous with frank “flat” mucosa and this should spur the quest for an understanding of the mechanisms underlying coeliac lesion. It is difficult to understand why gluten ingestion in these patients causes an uneven pattern of mucosal modification. Our data, demonstrating the presence of signs of local immune activation, indicate that immunological activation may not always lead to mucosal damage. Most remarkably, even fragments with normal histology produced detectable levels of EMA, indicating disease specific immune activation. These data suggest that EMA by themselves do not induce the histological modification observed in canonical flat mucosa, as recently suggested by Biagi and colleagues, although we cannot exclude the fact that EMA in combination with other factors may contribute to disease manifestation.

Our study also indicates that there is a hierarchical structure in even the epithelial element as reduction of apoptosis did not lead to a decrease in Ki67 expression. Thus it appears that the proliferative phase is not a consequence of induction of death but rather that proliferation precedes apoptosis. Elucidation of the factors controlling expression of FAS/apoptosis and induction of Ki67 could shed further light on the evolution of this pathology. Study of these patchy lesions may be the ideal tool to define the distinction between immunological activation and mucosal damage/remodelling, as immunological activation is a constant feature of untreated CD while mucosal damage/remodelling is not. The financial support of Telethon-Italy (Grant No E762), the Wellcome Trust (UK), Associazione Italiana Celiachia (Italy) European Community grants QLK1-CT-1999-00037, and BMH4-98-3703 is gratefully acknowledged. The Kennedy Institute of Rheumatology is supported by the Arthritis and Rheumatism Council UK.
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