An immunodominant DQ8 restricted gliadin peptide activates small intestinal immune response in in vitro cultured mucosa from HLA-DQ8 positive but not HLA-DQ8 negative coeliac patients

G Mazzarella, M Maglio, F Paparo, G Nardone, R Stefanile, L Greco, Y van de Wal, Y Kooy, F Koning, S Auricchio, R Troncone

Background: Studies on intestinal T cell clones from the mucosa of patients with coeliac disease have led to the identification of immunogenic gliadin epitopes. One is HLA-DQ8 restricted, its recognition by T cells being increased by introduction of negatively charged residues operated by tissue transglutaminase.

Aim: To test HLA-DQ8 restricted epitope in both native (QYPSGQGGSQPSQQNPQA) and deamidated (QYPSEGSGFQPSQGENPQA) forms in an organ culture system of treated coeliac mucosa from HLA-DQ8 positive and HLA-DQ8 negative patients.

Patients and methods: Jejunal biopsies obtained from 10 patients with coeliac disease (six HLA-DQ8 positive and four HLA-DQ8 negative) were cultured in vitro with a peptic-tryptic digest (PT) of gliadin, or with the native (peptide A) or deamidated (peptide B) peptide. Intraepithelial CD3+ and lamina propria total CD25+ cells were counted, lamina propria intercellular adhesion molecule 1 (ICAM-1) expression was evaluated, as well as that of Fas molecules on epithelial cells.

Results: In HLA-DQ8 positive, but not in HLA-DQ8 negative, coeliaics the density of intraepithelial CD3+ cells, lamina propria total CD25+ and CD3+CD25+ cells were significantly increased in biopsies cultured with PT, peptide A, or peptide B compared with biopsies cultured in medium alone.

Conclusion: These data show that the DQ8 restricted gliadin peptide is immunogenic only in the intestinal mucosa of HLA-DQ8 positive coeliac patients in both native and deamidated forms.

Celiac disease (CD) is one of the most thoroughly investigated and characterised of the many forms of food intolerance. It is defined as a permanent intolerance to wheat gliadins and related prolamine proteins, which provide enteropathy in genetically susceptible individuals.1 Architectural changes, phenotype of lamina propria mononuclear cells, and pattern of cytokines indicate a predominance of the Th1 response.1 Susceptibility to CD is determined to a significant extent by genetic factors; a great part of the genetic susceptibility maps to the HLA region on chromosome 6 as approximately 95% of CD patients carry an almost identical HLA DQ2 heterodimer (DQA1*0501 DQB1*0201). In most of those individuals negative for this haplotype the disease is associated with the class II alleles DQA1*0301 and DQB1*0302 that encode a DQ8 molecule which is linked to DRA4 haplotypes.5

Recent studies on intestinal T cell clones from the CD mucosa have led to the identification of a few immunogenic gliadin epitopes.3–5 Three are clearly HLA-DQ2 restricted; one is HLA-DQ8 restricted and its recognition by T cells is increased by introduction of negatively charged residues operated by tissue transglutaminase.4

Organ culture of the small intestine is a valuable model to study the immunological events occurring in the coeliac mucosa following contact with gliadin peptides; in fact, it has been shown that the in vitro gliadin challenge system reproduces many features of the mucosal immune response which occur in the established coeliac lesion.6 It has been proposed that gliadin challenge may initiate two parallel pathways, one of which leads to T cell activation and another that precedes it. Epithelial cells overexpress HLA-DR molecules within two hours, and in a second stage, T lymphocytes become fully activated; moreover, T lymphocytes migrate in the upper mucosal layers; T lymphocytes that migrate in the higher lamina propria compartments are mainly CD4+ and show markers of activation; and migrating intraepithelial lymphocytes are CD8+ and do not express these markers. The organ culture system has already been used to characterise gliadin epitopes able to activate mucosal cell mediated immunity.7

The aim of this study was to test the HLA-DQ8+ restricted gliadin epitope, in both native and deamidated forms, in an organ culture system.

MATERIALS AND METHODS

Patients and HLA typing

Ten patients with CD (four male and six female; median age 24 years (range 13–45)), diagnosed on the basis of revised ESPGHAN criteria, were recruited. They were on a gluten free diet for at least two years; they showed normal small intestinal histology and negative serology (antigliadin and antigliadin antibodies).

Molecular typing was performed as previously described.9 Briefly, DNA extraction was carried out according to a rapid
Table 1 HLA DQ molecular typing of subjects investigated in this study

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Peptides

A trypsin tryptic digest (PT) of gladin (variety S Pastore, gift of Dr De Vincenzi, Istituto Superiore di Sanita’, Rome, Italy) was used as a positive control. Peptide A corresponds to the residues 203–220 (sequence QYPSSGGSFQPSQNPQA of gda09, SwissProt P18573). Its deamidated form (peptide B) presents two residues of glutamine converted to glutamic acid (peptideB1m g / m l )peptide. The dishes were placed in a sterile nitrogen, and stored at 58°C until cryosectioning. The reaction product was developed by stirring continuously for three minutes with 20 mg of naphthol-AS-bisphosphate in 0.5 ml N-N-dimethylformamide added to 0.2 ml of sodium nitrite (Sigma Chemical Company, St Louis, Missouri, USA ), 0.08 ml of New Fucsin (Merck, West Point, Pennsylvania, USA), 40 ml of PBS, pH 8.7, and 17.5 mg of Levamisole (Sigma). Omission of the primary antibody served to control against non-specific antibody binding. The sections were finally stained with Mayer’s haematoxylin and mounted. Non-immune mouse immunoglobulins at the same isotype were used as the primary antibody for control of specificity. In experiments to detect activated T cells (CD3+CD25+), immunofluorescence with confocal microscopy was used because it allows better resolution of three colour procedures. Cryosections were fixed in acetone and incubated for one hour at room temperature with a mixture of mouse monoclonal antibodies anti-CD25 human (Dako, 1:25) and rabbit polyclonal anti-CD3 human (Dako, 1:100), followed by a mixture of horse antimouse FITC conjugated (1:200, Vector Laboratories, California, USA) and swine antirabbit TRITC conjugated (Dako, 1:300). These antibody were applied for 45 minutes in the dark followed by incubation (30 minutes) with ToPro-3 (Molecular Probes Leiden, the Netherlands) for counterstaining the nuclei. Finally, sections were mounted in phosphate buffered saline (PBS):glycerol (1:1). The primary and secondary antibodies were diluted appropriately in PBS containing bovine serum albumin at 1%, and all incubations took place at room temperature with intervening 10 minute PBS rinses.

CD3+ and CD25+ immunofluorescence cells were imaged with a Leica SP confocal microscope (Germany) using a 40x Plan-Neofluar oil immersion objective. The illumination source was from three single lasers: 488 nm line from argon ion laser, 543 nm line from HeNe ion laser, and 633 nm line from HeNe ion laser.

Morphometric analysis

The density of cells expressing CD3 in the intraepithelial compartment was determined by counting the number of stained cells as a percentage of 100 enterocytes; the number of CD3+ and CD25+ double positive cells was determined by counting the number of double positive cells as a percentage of all CD3 positive cells.

Table 2 Monoclonal and polyclonal antibodies used in this study

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Processing of jejunal biopsies and in vitro organ culture

Four biopsy specimens from the proximal jejunum were obtained from each patient, and all specimens were placed immediately in ice chilled 0.15 mol/l sodium chloride and brought to the laboratory within 30 minutes. One specimen from each patient was appropriately oriented and embedded in OCT compound (Tissue Tek, Miles Laboratories, Elkhart, Indiana, USA), snap frozen in isopentane, cooled in liquid nitrogen, and stored at −70°C until cryosectioning. The remaining specimens were placed on a stainless steel mesh positioned over the central well of an organ culture dish (Becton Dickinson, New York, USA) with the mucosal surface of the biopsies uppermost. Culture took place as previously reported. Here, the first six will be referred to as DQ8 positive and the remaining four as DQ8 negative.

Staining technique

Cryostat sections (5 µm) were air dried at room temperature and fixed in acetone for 10 minutes. All sections were repeatedly washed at room temperature in Tris buffered saline (TBS), incubated in normal rabbit serum (1:100 in TBS for 30 minutes), and then stained according to the peroxidase-antiperoxidase (PAP) or alkaline phosphatase/antialkaline phosphatase (APAAP) method. They were individually tested with the monoclonal antibodies (table 2). After one hour of incubation at room temperature, sections were incubated for 30 minutes with rabbit antimouse serum (Dako, Milan, Italy) and for a further 30 minutes with APAAP complex to stain CD25 positive cells and intercellular adhesion molecule 1 (ICAM-1) positive cells, or with PAP complex (Dako) to stain CD3 and CD95 (Fas) positive cells. Slides were washed in TBS for 10 minutes following each antibody incubation and were developed by a final incubation of five minutes with 2 amino-9 ethyl-carbazole (Sigma) when the PAP complex was used; when the APAAP complex was used instead, the immune reaction product was developed by stirring continuously for three minutes with 20 mg of naphthol-AS-bisphosphate in 0.5 ml N-N-dimethylformamide added to 0.2 ml of sodium nitrite (Sigma Chemical Company, St Louis, Missouri, USA ), 0.08 ml of New Fucsin (Merck, West Point, Pennsylvania, USA), 40 ml of TBS, pH 8.7, and 17.5 mg of Levamisole (Sigma). Omission of the primary antibody served to control against non-specific antibody binding. The sections were finally stained with Mayer’s haematoxylin and mounted. Non-immune mouse immunoglobulins at the same isotype were used as the primary antibody for control of specificity. In experiments to detect activated T cells (CD3–CD25+), immunofluorescence with confocal microscopy was used because it allows better resolution of three colour procedures. Cryosections were fixed in acetone and incubated for one hour at room temperature with a mixture of mouse monoclonal antibodies anti-CD25 human (Dako, 1:25) and rabbit polyclonal anti-CD3 human (Dako, 1:100), followed by a mixture of horse antimouse FITC conjugated (1:200, Vector Laboratories, California, USA) and swine antirabbit TRITC conjugated (Dako, 1:300). These antibody were applied for 45 minutes in the dark followed by incubation (30 minutes) with ToPro-3 (Molecular Probes Leiden, the Netherlands) for counterstaining the nuclei. Finally, sections were mounted in phosphate buffered saline (PBS):glycerol (1:1). The primary and secondary antibodies were diluted appropriately in PBS containing bovine serum albumin at 1%, and all incubations took place at room temperature with intervening 10 minute PBS rinses.

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lamina propria mononuclear cells (LPMNC) was evaluated within a total area of 1 mm\(^2\) of lamina propria using a microscope with a calibrated ocular aligned parallel to the muscularis mucosae. Lamina propria CD3\(^+\)CD25\(^+\) cells were counted as a percentage of the total CD3\(^+\) cells; at least 500 CD3\(^+\) cells were evaluated in each sample. Staining of epithelial cells by anti-CD95 and LPMNC by anti-ICAM-1 was evaluated in terms of staining intensity and graded on an arbitrary scale of weak staining (\(+=1\)), moderate staining (++\(=2\)), or strong staining (++\(+)=3\)). Counts were independently analysed in a blinded manner by two observers.

**Statistical analysis**

The Student’s two tailed \(t\) test was used to compare specimens exposed to gliadin or synthetic peptides with those exposed to medium alone for the same incubation time. Non-parametric tests (Wilcoxon two tailed) were also applied and the results are concordant with those obtained using parametric tests.

**RESULTS**

**Effects of in vitro HLA-DQ8 restricted peptide challenge on T cell activation**

For both HLA-DQ8\(^+\) and HLA-DQ8\(^-\) coeliac patients, in jejunal biopsies cultured in the presence of a PT digest of gliadin, the number of cells/mm\(^2\) expressing the interleukin 2 receptor (CD25 cells) (mean (SD) 121 (40) and 113 (15), respectively) was significantly (\(p<0.001\)) higher than in those cultured in medium alone (28 (6) and 26 (8), respectively) (fig 1). However, only in HLA-DQ8\(^+\) coelities was the number of CD25\(^+\) cells in specimens cultured in the presence of native peptide (139 (22)) or deamidated peptide (124 (21)) significantly higher (\(p<0.001\)) than in those cultured in medium alone. In fact, no significant differences were noted in HLA-DQ8\(^-\) coeliac patients between biopsy specimens cultured with native peptide (37 (2)) or deamidated peptide (38 (10)) and those cultured in only medium (26 (8)) (fig 1). In further experiments, we examined the increase in CD3\(^-\)CD25\(^-\) cells; in fact, both HLA-DQ8\(^-\) and HLA-DQ8\(^+\) coeliac patients challenged in vitro with the PT digest of gliadin showed a significant increase (\(p<0.01\)) in activated T cells; nevertheless, only HLA-DQ8\(^+\) coeliac patients showed a significant increase (\(p<0.05\)) of CD3\(^-\)CD25\(^-\) cells in response to native or deamidated peptide (table 3, fig 2).

Expression of ICAM-1 was also evaluated to find further evidence of activated mucosal cell mediated immunity. ICAM-1 was increased significantly in all cells and vessels of biopsy specimens from HLA-DQ8\(^+\) (\(p<0.001\)) and HLA-DQ8\(^-\) (\(p<0.01\)) coeliac patients cultured for 24 hours with PT gliadin compared with those cultured with medium alone; in contrast, only in coeliac HLA-DQ8\(^+\) was ICAM-1 expression...
Effects of in vitro HLA-DQ8 restricted peptide challenge on T cell infiltration

A significant increase in CD3 intraepithelial lymphocytes was seen when biopsies of HLA-DQ8 patients were cultured in the presence of native peptide (23 (9)) (fig 4, fig 2) and HLA-DQ8 coeliac patients cultured in vitro with PT gliadin, both in biopsy samples obtained from HLA-DQ8 (p<0.001) and HLA-DQ8 (p<0.01) coeliac patients compared with samples cultured in the presence of medium alone (fig 5). Also, significantly enhanced expression was seen when HLA-DQ8 coeliac biopsies were cultured in the presence of native peptide (p<0.001) or deamidated peptide (p<0.01) compared with those cultured in medium alone (fig 4, fig 5). In HLA-DQ8 coeliac patients, a significant (p<0.01) increase in CD3 intraepithelial lymphocytes was seen only in biopsy specimens cultured in the presence of gliadin (42 (12)). Significantly enhanced expression was seen when HLA-DQ8 coeliac patients were cultured in the presence of the DQ8 restricted peptides (both native 26 (8) and deamidated 25 (4)) compared with those cultured in medium alone (21 (8)) (fig 4).

DISCUSSION

Most evidence suggests that a decisive role in the pathogenesis of coeliac disease is played by HLA-DQ2 and/or DQ8 restricted peptides (both native and deamidated).
Immunodominant DQ8 restricted gliadin peptide in coeliac patients

Figure 6  Fas expression in the epithelium of jejunal mucosa from a DQ8 coeliac patient cultured in vitro with medium only (A) or with native peptide (peptide A) (B). In the latter, intense staining is detected in almost all of the epithelial cells; expression is particularly evident in basolateral membranes of enterocytes. A similar pattern was observed when the mucosa was cultured with the deamidated peptide or with the peptic-tryptic digest of gliadin. Original magnification ×100.

gluten specific T cells at the site of the lesion in the gut. None the less, because of the considerable biochemical complexity of gluten proteins, there is still uncertainty about the amino acid sequences which are responsible for mucosal damage. Several tools have been used over the years to identify “toxic” sequences, probably investigating different properties of gliadin peptides not always related to T cell recognition. Most recently, considerable progress in the identification of T cell stimulatory gluten derived peptides has been rendered possible by the isolation of gliadin specific T cell clones from the mucosa of treated and untreated patients, and by the implementation of modern analytical tools. This work has proved that deamidation can enhance T cell reactivity to such peptides. To date, five unique epitopes of gliadin that are recognised by T cells have been identified, three restricted by HLA-DQ2 and two by HLA-DQ8 molecules, but many more are predicted to be immunogenic. Reactivity of all HLA-DQ2 restricted epitopes and one of the HLA-DQ8 restricted peptides is significantly enhanced by deamidation of specific glutamine residues to glutamic acid. This does not seem to be true for the remaining HLA-DQ8 determinant which belongs to a glutenin molecule.

We aimed to transfer these observations to a more complex system represented by the in vitro organ culture of coeliac biopsies, a model already used in the past to test the biological properties of gliadin peptides and, more recently, their immunogenicity. Our data showed that the DQ8 restricted gliadin epitope, identified for its activity on T cell clones, was also active in the organ culture system of the treated coeliac mucosa. In this system, it significantly increased the density of lamina propria mononuclear cells and in particular T cells expressing the interleukin 2 receptor. The extent of the latter phenomenon, which is accompanied by a redistribution of these cells, recruited in the subepithelial compartment, is of similar amplitude to that observed in cultures performed in the presence of the whole gliadin PT digest. The most likely explanation is that the peptide used is certainly immunodominant but it is also possible that activation of T cell clones specific for that peptide is followed by a bystander activation of other mononuclear cells; in fact, in our culture system, following gliadin stimulation, the dominating CD25 subset was represented by CD3-CD25 cells, the latter being presumably activated macrophages.

We have also observed significant enhancement of expression of ICAM-1 in those fragments cultured in the presence of the peptide. The latter phenomenon has already been described in treated coeliac biopsies cultured with a gliadin digest and is attributed to the strong γ-interferon induction which is one of the main features of this in vitro response. Moreover, two other phenomena have been observed in our biopsies cultured with the peptide: increased intraepithelial density of CD3 cells and enhanced expression of Fas molecules on epithelial cells. The first of such observations is a phenomenon already shown to occur in vitro in response to gliadin; as it was not affected by the presence of CTLA4-Ig, a molecule interfering with costimulation of T cells, its T cell dependence has been questioned, and a pathway parallel to that involving mucosal T cells has been hypothesised. Similar observations have been made for induction of Fas for which a direct effect on epithelial cells (non-T cell mediated) has been suggested. Yet, it is still possible that CD3 epithelial infiltration is a consequence of lamina propria T cell activation. In fact, in explants of human fetal gut, small intestine T cell activation by anti-CD3 antibodies induced a significant increase in intraepithelial lymphocytes, suggesting that it might be a consequence of lamina propria T cell activation. More difficult to explain is Fas expression on epithelial cells as there is no previous evidence suggesting the influence of T cell activation on such phenomenon, which in our experience seems to be DQ restricted. The relatively large amount of peptide used in the system must also be considered in interpreting the results.

None of these phenomena was observed in HLA-DQ8 negative subjects. This clear HLA-DQ restriction of peptide activity has strong implications not only for our understanding of the sequence of events triggered by gliadin in the coeliac mucosa; in fact, for the first time gliadin peptides have been shown to display different biological activities in different subsets of coeliac patients. It is still unclear if the response of coeliac patients is directed against a few or multiple peptides, but our observations raise the issue of their immunogenicity, depending on the genetic make up of the patient. For the peptide that we investigated, it is likely to be immunodominant as all six HLA-DQ8 coeliac mucosae tested showed a clear response. We have also observed significant enhancement of expression of ICAM-1 in those fragments cultured in the presence of the peptide. The latter phenomenon has already been described in treated coeliac biopsies cultured with a gliadin digest and is attributed to the strong γ-interferon induction which is one of the main features of this in vitro response. Moreover, two other phenomena have been observed in our biopsies cultured with the peptide: increased intraepithelial density of CD3 cells and enhanced expression of Fas molecules on epithelial cells. The first of such observations is a phenomenon already shown to occur in vitro in response to gliadin; as it was not affected by the presence of CTLA4-Ig, a molecule interfering with costimulation of T cells, its T cell dependence has been questioned, and a pathway parallel to that involving mucosal T cells has been hypothesised. Similar observations have been made for induction of Fas for which a direct effect on epithelial cells (non-T cell mediated) has been suggested. Yet, it is still possible that CD3 epithelial infiltration is a consequence of lamina propria T cell activation. In fact, in explants of human fetal gut, small intestine T cell activation by anti-CD3 antibodies induced a significant increase in intraepithelial lymphocytes, suggesting that it might be a consequence of lamina propria T cell activation. More difficult to explain is Fas expression on epithelial cells as there is no previous evidence suggesting the influence of T cell activation on such phenomenon, which in our experience seems to be DQ restricted. The relatively large amount of peptide used in the system must also be considered in interpreting the results.

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In conclusion, we have evidence that a peptide identified and characterised for its activity on T cell clones isolated from
coeliac mucosae is also active, in a HLA-DQ restricted manner, in an in vitro organ culture system, causing T cell activation, intraepithelial migration of T cells, and preapoptotic phenomena. The organ culture system is confirmed as very useful in determining the biological activity of gliadin peptides, before the ultimate proof of their toxicity is provided by challenge studies.

ACKNOWLEDGEMENTS
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