Autoantibodies to gut hormone secreting cells as markers of peptide deficiency

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SUMMARY Autoantibodies reacting with endocrine cells in the gastrointestinal mucosa were found by indirect immunofluorescence in 22 out of 268 sera (8.2%) obtained from patients with coeliac disease, Crohn's disease, ulcerative colitis, irritable bowel syndrome, and from subjects without bowel disease. A double immunofluorescence technique showed that the autoantibodies reacted with cells secreting gastric inhibitory polypeptide (glucose dependent insulino tropic polypeptide, GIP), secretin, somatostatin or enteroglucagon. Most sera contained antibodies against more than one cell type. Neither the presence of a particular antibody nor the pattern of antibody combinations appeared to be specific for any diagnostic category. The mean plasma GIP concentrations, however, both fasting and two hours after a test meal, were significantly lower in subjects with GIP cell autoantibodies. Thus gut hormone cell autoantibodies may be markers of impaired hormone secretion.

Organ specific circulating autoantibodies which can be detected by indirect immunofluorescence are frequently associated with disturbed endocrine function – for example, autoimmune thyroid disease, Addison's disease, type I insulin dependent diabetes, etc. Recent advances in immunocytochemistry have established the existence of an extensive endocrine system within the mucosa of the gastrointestinal tract. The possibility of humoral autoimmunity directed against this endocrine tissue was suggested by reports of autoantibodies against a number of gut hormone cells in diabetes mellitus (pancreatic glucagon and somatostatin cells), atrophic gastritis (gastrin cells), and coeliac disease (duodenal GIP and secretin cells) but no systematic search has been reported in other chronic bowel disorders. Autoantibodies against peptide secreting cells throughout the gut were therefore sought in sera from patients with a variety of intestinal disorders and in subjects without evidence of bowel disease.

Methods

SUBJECTS

Serum samples were tested from a total of 268 subjects (115 men, 153 women). The diagnoses in these were established by accepted clinical, radiological, or histological criteria and included coeliac disease (43 patients), dermatitis herpetiformis (24 patients, 10 of whom also had coeliac disease), Crohn's disease (47), ulcerative colitis (65), and irritable bowel syndrome (44). The remaining 45 subjects included 30 patients without evidence of past or present bowel disorder and 15 healthy members of hospital staff. Follow up serum samples after intervals of three to 18 months were obtained from 20 subjects in whom antibodies to gut endocrine cells were detected in the initial samples.

TISSUE SUBSTRATES

'Normal' intestinal mucosa was obtained at operation from patients of blood group O undergoing surgery because of pathology, usually neoplasia, in adjacent bowel. Samples of duodenum, jejunum, ileum, and colon from several different donors were used (Table 1). The intestinal
mucosa was immediately separated from the under-
lying muscle, rolled up, snap frozen in melting
isopentane, and stored in liquid nitrogen. Human
thyroid, stomach and pancreas, and rat liver and
kidney were also used in identifying other organ
and non-organ specific autoantibodies.

IMMUNOFLUORESCENCE METHODS

(a) Detection of autoantibodies Test sera were
applied for 60 minutes, without being previously
diluted, to tissue sections cut at 2–4 μm. Sections
were washed for 20 minutes in buffered saline and
sheep fluorescein (FITC) conjugated antihuman
whole immunoglobulin, anti-IgG, -IgA, and -IgM
(Wellcome Laboratories), or sheep antihuman C₃
(Nordic Laboratories) together with fresh normal
human serum as a source of complement were
applied. Antibody titres were determined by serial
doubling dilutions. All sera were tested on mucosa
from at least two different donors.

(b) Identification of gut hormone cells A four-layer
immuno-fluorescence technique was used as
previously described. After the sequence described
in (a) specific rabbit antisera against porcine GIP,
glucagon and vasoactive intestinal polypeptide
(VIP), or synthetic secretin and somatostatin
(kindly provided by various laboratories as detailed
previously)³ were applied followed by goat
rhodamine (TRITC) conjugated antirabbit immuno-
globulin (Wellcome).

SPECIFICITY OF AUTOANTIBODIES

1 Absorption with specific hormone In order to
be certain that the fluorescence was because of
antibodies directed against components of the cells
other than the hormones themselves, two sera,
containing antibodies against GIP cells but not
against other cell types, were absorbed at 4°C
overnight with an excess of pure GIP, to give a final
serum concentration of 250 000 pmols of GIP per
litre. Control sera were absorbed with buffered
saline alone. An identical series of experiments was
 carried out with GIP hormone antiserum at a
concentration of four doubling dilutions below
endpoint.

2 Inactivation of complement components In
order to exclude non-specific complement-
mediated binding of immunoglobulins to endocrine
cells the following two procedures were undertaken
before applying sera to frozen sections as described
above: (i) Firstly, aliquots of 20 sera containing
putative autoantibodies were absorbed at 4°C
overnight with an excess of antihuman C₁q (200 μl
serum, 20 μl anti-C₁q, DAKO, Mercia Borcades Ltd
batch No 129A, titre 11 280). Control sera were
absorbed with 20 μl of buffered saline. (ii) Secondly,
to inactivate heat labile components of complement,
aliquots of 20 positive sera were heated to 56°C for
one hour.

METABOLIC RESPONSES

As subjects with GIP cell autoantibodies were the
most numerous it was decided to assess their
metabolic responses to a test meal. Eleven subjects
with GIP cell autoantibodies were compared with 11
subjects without these antibodies. There were nine
women and two men in each group matched for age
and disease (Table 2). After a fasting basal blood
collecting 2225 KJ 'standard breakfast' was given
and further blood was obtained after 30 and 120
minutes. Blood for hormone assay was collected in
heparinised tubes containing aprotinin (400
kallikrein inactivating units per ml of blood). The
plasma was separated within 15 minutes and stored
at −20°C until assayed without knowledge of the
antibody results. Simultaneous samples were
analysed for whole blood glucose concentration
using glucose oxidase. Hormone profiles were
compared in the matched pairs using a two-tailed
paired t test.

Table 2 Numbers of sera tested in each diagnostic category and numbers containing autoantibodies

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<th>Numbers of sera tested</th>
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<td>Total</td>
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<table>
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<td>Without evident bowel disease</td>
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Figures in parentheses show the number of pairs of patients who underwent metabolic studies.
**Autoantibodies to gut hormone secreting cells as markers of peptide deficiency**

**HORMONE RADIOIMMUNOASSAY**

Plasma GIP and insulin were measured by conventional radioimmunoassays. The GIP antiserum was raised to pure natural porcine material and fully detected both big and little human GIP though showing no cross reaction with other peptides.

**Results**

**IMMUNOFLUORESCENCE**

Twenty-two of the 268 sera (8.2%) contained antibodies which reacted with mucosal cells in the duodenum and jejenum having a distribution and morphology suggestive of endocrine cells. Twenty of these sera were tested against all the tissue substrates collected (Table 1) and gave positive results with the appropriate mucosa in all cases. The remaining two sera were tested on duodenum from two donors only, both giving identical results. Three of these 22 sera also reacted with cells in the colonic mucosa. No fluorescence reactions were obtained on ileal mucosa; all the sera were tested on the ileal mucosa from two donors and many of them on an additional three substrates. The antibodies in all the 22 positive sera were of IgG class and there was an additional IgM component in eight of them. No reactions were obtained using anti-IgA conjugate. Results were unaffected by the previous heating of serum to 56°C for one hour or by overnight absorption with anti-C1q. Using anti-C3 conjugate all of 18 sera tested were shown to fix complement after the addition of fresh normal human serum. Antibody titres using polyvalent anti-human immunoglobulin ranged from 1:4 to 1:32. Further serum samples obtained from 20 of the subjects three to 18 months later produced identical fluorescent patterns and titres. These antibodies did not correlate with any of the clinical diagnoses included in this study and occurred in each of the groups tested (Table 2). There was no discernible difference in disease activity or duration of disease in patients with gut endocrine cell antibodies compared with patients of the same diagnostic category but without antibodies. Subjects without bowel disease but with gut hormone cell auto-antibodies included three healthy volunteers, one patient with chronic bronchitis, and one patient with mild dementia.

Four layer immunofluorescence confirmed that these antibodies were reacting with mucosal endocrine cells. Most of the sera contained antibodies against more than one cell type (Table 3). The pattern of the reacting cells among the 22 sera again appeared to be independent of diagnostic category. Overnight absorption with pure GIP failed to affect the fluorescence produced with two sera containing GIP cell autoantibodies while abolishing the fluorescence obtained using a GIP hormone antiserum on frozen sections.

**METABOLIC RESPONSES TO A STANDARD MEAL**

Plasma GIP levels in subjects with and without GIP cell autoantibodies are shown in Fig. 1. The mean basal plasma level was significantly lower in antibody positive subjects (6.82±0.71 pmol/l) than in antibody negative subjects (10.82±1.02 pmol/l; p<0.02); a significantly lower mean value was also obtained 120 minutes postprandially in the subjects with antibodies (21.10±2.14 pmol/l, compared with 26.64±1.9 pmol/l; p<0.02). Values 30 minutes postprandially were not significantly different between the two groups.

Plasma insulin responses did not differ in the two groups (Fig. 2). Whole blood glucose levels were lower in the fasting samples for patients with GIP cell antibodies than in those without but postprandial values were similar in the two groups (Fig. 3). No fasting blood sugar value exceeded 6 mmol/l.

**Discussion**

Despite earlier reports that autoantibodies occur against duodenal GIP and secretin cells in coeliac disease and tropical sprue, the significance of such antibodies has not been determined. No previous systematic search for autoantibodies reacting with peptide secreting cells throughout the gut in different gastrointestinal diseases has been reported. This study confirms the existence of a system of humoral autoimmune directed against several different gut peptide secreting cells. The
Table 3 Clinical details of 22 subjects with gut endocrine cell autoantibodies

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* Note that when more than one endocrine cell antibody occurs in a single serum only the overall titre can be assessed.

Autoantibodies to gut hormone secreting cells as markers of peptide deficiency

antibodies are not directed against the secreted hormone as shown by the failure to abolish GIP cell fluorescence after absorption with pure GIP. In common with other endocrine autoantibodies, therefore, they appear to react with structural cytoplasmic components.

It has been suggested that hormone secreting cells of the gastrointestinal mucosa may bind immunoglobulins in a non-specific manner, mediated via the C₁q fraction of complement, producing fluorescence on fixed sections. This does not explain the results in the present study because heating the sera to 56°C, or absorption with anti-C₁q, failed to abolish fluorescence on frozen sections.

As it was possible to observe varying combinations of reacting cell types using sera from different patients it is likely that distinct antigens are present in each of the different hormone secreting cells. A similar observation has been made in relation to pancreatic islet and pituitary cells.

The finding that the mean plasma GIP concentrations are lower in the group of patients with GIP cell antibodies suggests that the antibodies may be markers for a pathological process affecting those cells. The assessment of any possible cytotoxic effect of these autoantibodies must await the development of techniques for the accurate separation and quantification of gut peptide cells. By determining metabolic responses in subjects with isolated peptide cell abnormalities, however, it is likely that information at least comparable with that of hormone infusion experiments will be obtained. Likewise, future studies of gut hormone responses should take account of the existence of this autoimmune system.

The concept of GIP as a glucose dependent insulino tropic hormone must be considered. It has been suggested that GIP may play a major part in the enteroinsular response to ingested glucose and be important in diabetes mellitus, particularly of the type II non-insulin dependent variety. Fasting blood glucose levels, however, were below the accepted value for a diagnosis of diabetes mellitus in all subjects. It is of interest that among these non-diabetic patients with GIP cell antibodies there was a decreased GIP response to a test meal while the glucose responses and insulin levels did not differ from the control group with no such antibodies. Nevertheless, the possibility remains that the presence of GIP cell autoantibodies may be important in patients with established diabetes mellitus.

The present data do not suggest a specific association between the presence of circulating gut endocrine cell antibodies and any of the gastroenterological disorders investigated so far. If the actions of the peptides, however, are mainly at a local level on adjacent cells - that is, paracrine - then these antibodies might be expected to provide new insight into the influence of individual cells upon their neighbours. As 'gut hormones' are widely distributed throughout the body - for example, as neurotransmitters - the possibility remains that the antibodies may reflect pathology in another organ system entirely. Further work is indicated, therefore, to determine their prevalence in other conditions.

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References