Case report

Alpha heavy chain disease lacking secretory alpha chain, with cobblestone appearance of the small intestine and duodenal ulcer demonstrated by endoscopy

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SUMMARY Ultrastructural and immunohistochemical studies of the small intestine are described in a Japanese patient with alpha heavy chain disease who had a history of colonic ulcers. Endoscopic examinations revealed multiple gastric erosion, duodenal ulcer, and a thickened, cobblestone-like pattern composed of small nodules in the duodenum and jejunum, which was similar in appearance to Crohn's disease. An electron microscopic study showed that the numerous, infiltrated cells in the jejunal lamina propria were matured plasma cells with atypical structure of the organelles. These plasma cells had alpha heavy chain protein devoid of light chain. Although secretory component was demonstrated normally in the epithelial cells by immunofluorescent methods, no association of this component with alpha heavy chain protein could be observed in any of the plasma and epithelial cells of this case. These facts suggest the absence of secretory alpha chain or secretory IgA, and a deficiency of the mucosal secretory immune system in this patient.

Since the first description of alpha heavy chain disease more than 100 cases have been reported. This disorder appears to be a condition primarily affecting the secretory IgA system of the digestive tract. For studies on the secretory immune system, alpha heavy chain disease represents an appropriate natural model. Whether the binding of secretory component with alpha heavy chain protein in the intestine is present or not in this disorder, however, has not yet been clarified.

The disorder has been observed predominantly in the Mediterranean area and the Middle East. Although a few cases have been reported in other Eastern countries, only two cases have so far been described in Japan.2 3

The purpose of this paper is to report a third case of alpha heavy chain disease in Japan, associated with multiple gastric erosion and duodenal ulcer. Furthermore, this case was investigated by electron microscopic and immunohistochemical methods in order to clarify the localisation of secretory component in the small intestine.

Case report

A 39-year-old Japanese male developed occasional diarrhoea associated with lower abdominal pain in March 1975. The next year he suffered from severe diarrhoea. At that time, the possibility of Crohn's disease was considered, as multiple ulcers with a cobblestone-like appearance were demonstrated in the duodenum and ileocecal regions by barium enema radiographs and endoscopic examinations. In July 1977, the patient was admitted to Ashikaga Red Cross Hospital because of severe diarrhoea, abdominal pain, and fever.

Physical examination revealed nothing noteworthy except for abdominal distension. Laboratory
investigations gave a haemoglobin level of 6-3 g/dl, white blood cell count of 5100/mm³ (neutrophils 68%, lymphocytes 30%, monocytes 2%, eosinophils 0%, and basophils 0%) and platelets 21×10⁶/mm³. Serum electrolytes were within the normal range. The total protein was 4-4 g/dl (albumin 41-5%), total cholesterol 72 mg/dl, LDH 209 W units, GOT 18 K units, GPT 24 K units, and alkaline phosphatase 10-2 KA units.

**Protein Studies**
Electrophoresis and immunoelectrophoresis were performed by standard methods. Serum immunoglobulins were estimated using radial immunodiffusion methods. Immunoexclusion methods were performed in 1-2% agar+agarose mixed gel, incorporated with 10% anti-lambda and kappa serum (Dekopatts, 500 titre). After electrophoresis for two hours at 2-5 mA/cm, anti-IgA (alpha chain) and anti-secretory component were loaded and allowed to diffuse overnight.

**Endoscopic Examinations**
An Olympus small intestine fibroscope (Olympus SIF-B) was used for observations of the stomach, duodenum, and jejunum. The fibroscope was inserted into the jejunum 30 cm distal to the ligament of Treitz.

**Immunohistochemical Studies**
Biopsy specimens from the jejunal mucosa through the Olympus SIF-B were fixed in 95% ethanol at 4°C overnight. After dehydration for three days in absolute ethanol, the tissues were embedded in paraffin and then cut into 4μ sections. Immunofluorescent studies were performed on these sections, according to the direct methods using monospecific antisera against alpha, gamma, mu, epsilon, kappa, and lambda human chains conjugated to fluorescein isothiocyanate (Behringerwerke, West Germany). Indirect methods using rabbit anti-human secretory component and goat anti-rabbit IgG conjugated with fluorescein isothiocyanate, were used to observe the localisation of secretory component. The specificity was assessed by the blocking test.

**Electron Microscopic Examinations**
The biopsied specimens from the jejunal mucosa were fixed with 2% glutaraldehyde in phosphate buffer and post-fixed with 1% osmic acid. After dehydration in graded solutions of ethanol, the materials were embedded in Epon 812. Thin sections were stained with uranyl acetate and lead citrate, and observed under a Hitachi HS-9 electron microscope.

**Results**

**Protein Studies**
The serum protein fraction showed a slight decrease in the gamma globulin fraction. The most striking abnormality was a broad rise in the alpha 2 and beta mobility range. Immunoelectrophoretic analysis of the patient's serum demonstrated a heavy and broadly dispersed precipitin arc which was developed with anti-IgA serum. The same arc was discernible in the immunoelectrophoretic pattern of the patient's serum developed with anti-Ig (G+A+M). Precipitin arcs corresponding to the IgA components were not found when the serum was developed with anti-Bence Jones kappa and lambda serum. Investigations by the immunoselection plate method were performed on the patient's serum. The serum did not react to anti-lambda and kappa serum, and showed the same bands as those seen in the immunoelectrophoresis (Fig. 1).

**Endoscopic Examinations**
The gastric mucosa showed multiple erosion with atrophic gastritis, and active ulcer was observed on the anterior wall of the duodenal bulb (Fig. 2). Through the second portion of the duodenum and jejunum, the mucosal surface consisted of nodules resembling cobblestones (Fig. 3). The size of each nodule, however, was smaller than that seen in Crohn's disease and the lesions were diffuse over the section of the intestine that was observed. Microscopically, biopsied specimens revealed villous atrophy with marked reduction in the height of the epithelial cells. The lamina propria was markedly infiltrated by plasmacytoid cells. These plasmacytoid cells showed infiltration into some parts of the submucosa.

**Immunohistochemical Studies**
The numerous cells infiltrating into the lamina propria exhibited a slight cytoplasmic fluorescence with anti-alpha chain serum (Fig. 4), and a few of them exhibited fluorescence with anti-kappa and lambda chain serum (Fig. 5). Gamma, mu, and epsilon positive cells were sparsely observed in the lamina propria. Moreover, the epithelial cells showed no fluorescence with anti-alpha chain serum, but clear fluorescence with anti-secretory component serum in the apical portions (Fig. 6). The localisation of the secretory component was predominantly in the crypt cells.

**Electron Microscopic Examinations**
Almost all of the infiltrating cells into the lamina propria revealed the mature pattern of plasma cells.
Fig. 1 Immunoselection plates showing the presence of alpha chain free of light chain. PS: patient serum; CS: control serum.

Fig. 2 Endoscopic features of the duodenal ulcer. Active ulcer with fold convergence is indicated.

Fig. 3 Endoscopic features of the jejunal mucosa. A cobblestone-like appearance with small nodules along the whole small intestine is observed.

Fig. 4 Alpha chain containing cells in the biopsied jejunal mucosa, using anti-alpha chain serum conjugated with FITC. Numerous alpha chain cells are seen in the lamina propria, but no alpha chain is localised in the epithelial cells. ×200 (original magnification).

Their nuclei were oval, with peripheral clumps of chromatin. Nucleoli were also abundant. The well-developed Golgi apparatus and dilatation of rough-surfaced endoplasmic reticulum, which suggested an increase of secretory activity, were observed in the cytoplasm (Fig. 7a). Some of the rough-surfaced endoplasmic reticulum was arranged in unusual whorl-like configurations (Fig. 7b).

Discussion

Alpha heavy chain disease is characterised by plasmacytic infiltration of the small intestine, the cells of which are considered to secrete an abnormal protein related to the alpha heavy chain of the IgA₁ subclass.⁴ The patients with alpha heavy chain disease described since 1968¹ have characteristically
been young and have come predominantly from the Middle East and Mediterranean areas. For this reason, it is important to report new cases outside these areas. In Japan, only three cases with alpha heavy chain disease, including our case, have been reported.\textsuperscript{2,3} Currently, the view is widely held that alpha heavy chain disease occurs as a reaction to chronic stimulation of the intestinal tract, because of the marked plasma cell infiltration and because the disease mainly occurs in areas of poor hygiene and with a high incidence of parasitic infiltration. Increase of epsilon chain cells, however, which has often been noted in cases of parasitic infection, was not observed in the present patient’s intestine.

Endoscopic examination of the upper gastrointestinal tract has never been reported previously, although there have been some descriptions of the colonic lesions of alpha heavy chain disease.\textsuperscript{5-8} Among reported cases whose stomach was studied by radiographic examination,\textsuperscript{9-11} none revealed gastric lesions, such as multiple erosion, except for one case in which polyposis was found in the stomach.\textsuperscript{12} An association with duodenal ulcer was reported in two cases.\textsuperscript{13,14} Studies on the stomach and duodenum in alpha heavy chain disease thus appear to be incomplete based on the literature (Table). The fact that multiple gastric erosion and duodenal ulcer were observed in the present patient...
suggests a breakdown of the protection mechanisms of the gastroduodenal mucosa in this disorder. Furthermore, the endoscopic findings such as the cobblestone-like appearance in the jejenum simulate those of Crohn's disease. Wang (1976)\textsuperscript{15} reported an English patient with alpha heavy chain disease in whom the possibility of Crohn's disease was considered on the basis of a barium meal study. More precise study on the diagnosis of Crohn's disease with high serum IgA values is thus warranted.

Biosynthetic studies performed on lymphoplasmacytic cells and tumour cells from the gut of patients with alpha heavy chain disease have revealed the synthesis of alpha heavy chain protein.\textsuperscript{16} Immunofluorescence studies of the gut performed in patients with the 'benign phase' have usually shown slight cytoplasmic fluorescence with anti-alpha chain antisera and no fluorescence with anti-light chain antisera.\textsuperscript{11} On the other hand, in patients with the 'malignant phase' the infiltrating cells have been completely negative with all antisera. Judged by the earlier reports (Table), the population of alpha chain positive cells has been variable, as the cases were not classified as being either benign or malignant.\textsuperscript{15, 17} In our case, which was of the benign phase as confirmed by the histopathological and electron microscopic examinations, most of the infiltrating cells were positive with anti-alpha chain antiserum, and almost negative with anti-light chain antiserum. From these findings, it is presumed that alpha chain protein is synthesised by infiltrating lymphoplasmacytic cells.

Secretory component in the epithelial cells of the jejenum from patients with alpha heavy chain disease has not been described previously (Table). Our patient's epithelial cells in the crypt contained secretory component in their apical portions, but did not possess alpha heavy chain in the same portions, as demonstrated by immunofluorescence studies. Moreover, the lamina propria consisted of alpha heavy chain positive cells. These findings suggest that secretory component and alpha heavy chain are produced separately in the small intestine, and that an association between the two did not occur in the present patient. This conflicts with the earliest report\textsuperscript{13} that the jejunal fluid from alpha heavy chain disease contained alpha heavy chain protein associated with secretory component, but agrees with another report\textsuperscript{20} that jejunal fluid from the patient did not contain secretory component (Table). These conflicting data remain unresolved. One possibility is that alpha heavy chain protein cannot possess the double Y configuration as in the structure of normal secretory IgA\textsuperscript{21} and that such structural alteration may cause difficulty for secretory component to bind alpha chain protein. Another possibility is that J chain might not be

\begin{table}
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\begin{tabular}{|c|c|c|c|c|}
\hline
\textbf{Authors} & \textbf{GI tract lesions except in small intestine} & \textbf{Authors} & \textbf{Immunohistochemical studies on intestinal mucosa} & \textbf{Salivary or jejunal fluids} \\
\hline
Rambaud\textsuperscript{13} & Duodenal ulcer & Seligmann\textsuperscript{19} & Variable & +
\textsuperscript{nd}\textsuperscript{*} & Rambaud\textsuperscript{13} \\
(1968) & & (1969) & nd & + & (1968) \\
Manousos\textsuperscript{9} & Nodular appearance & Doe\textsuperscript{20} & Variable & +
\textsuperscript{nd} & Stoop\textsuperscript{22} & Normal IgA \\
Plesnic\r\textsuperscript{5} & Numerous ulcerated & Manousos\textsuperscript{9} & Absent & +
\textsuperscript{nd} & Doe\textsuperscript{20} & (1972) \\
Shulmann\textsuperscript{6} & Multiple mucosal & Shulmann\textsuperscript{6} & Absent & +
\textsuperscript{nd} & Manousos\textsuperscript{9} & (1974) \\
Tabbane\textsuperscript{7} & Tumour involvement & Wang\textsuperscript{15} & Numerous & +
\textsuperscript{nd} & Plesnic\textsuperscript{5} & (1975) \\
Doe\textsuperscript{14} & Duodenal ulcer & Galian\textsuperscript{11} & Numerous & +
\textsuperscript{nd} & Shulmann\textsuperscript{6} & (1975) \\
Galian\textsuperscript{11} & Polypoid features & Haghshenas\textsuperscript{23} & Numerous & +
\textsuperscript{nd} & Haghshenas & (1977) \\
(1978) & in rectum & & & + & (1977) \\
Al-Bahrami\textsuperscript{8} & Extensive colonic & Gafter\textsuperscript{24} & & & \\
Cohen\textsuperscript{12} & Polypoid features & & & & \\
(1978) & in stomach and colon & & & & \\
Hibi\textsuperscript{17} & Multiple gastric & Hibi & Numerous & Present in the \\
duodenal ulcer & & & epithelial cells & & \\
\hline
\end{tabular}
\caption{Clinical and immunochemical studies of alpha heavy chain disease in literature}
\end{table}

* nd: not done.
produced in the plasmacytoid cells in this disorder. J chain is considered to be necessary for the binding of alpha heavy chain and secretory component. Unfortunately, we were unable to study J chain because of a shortage of purified anti-J chain antiserum.

In order to clarify the problem of binding, it is necessary to demonstrate secretory alpha chain localised in the intestinal epithelial cells. It is well established that IgA is synthesised by intestinal plasma cells and then binds to secretory component produced by epithelial cells in the normal intestine.25 26 Our findings suggest that secretory IgA or secretory alpha chain might not be produced in the intestine in the present patient. The absence of secretory alpha chain may cause a breakdown of the mucosal defence and render it easy for intestinal bacteria or viruses to invade the mucosa, although the role of alpha heavy chain protein devoid of light chain as antibody is unclear. This condition leads to a chance of intestinal tract stimulation and such weak mucosal defence mechanisms result in hyper-susceptibility to intestinal stimulants and overproduction of useless alpha heavy chain secreting cells.

References


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