Chronic gastritis and gastric ulcer

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EDITORIAL SYNOPSIS The authors focus attention on the possible role of chronic gastritis as a cause of gastric ulceration and bleeding resulting from decreased mucosal resistance.

Duodenal and gastric antral ulcers differ in certain respects from ulcers in the body of the stomach. Thus with duodenal and gastric antral ulcers there is a high gastric acidity, a large parietal cell mass, and a histologically normal gastric mucosa (Cox, 1952; Ball, 1961), whereas with gastric ulcers there is a normal, low, or even greatly reduced, gastric acid secretion (Levin, Kirsner, Palmer, and Butler, 1948; Morlock and Ratke, 1949; Ball, 1961; Baron, 1963), a reduced parietal cell mass, and often a histologically abnormal gastric mucosa with changes of gastritis (Magnus, 1952; Ball and James, 1961). Hence 'decreased mucosal resistance' rather than hyperacidity is claimed to be the important factor in the pathogenesis of gastric ulcer. Hence 'decreased mucosal resistance' rather than hyperacidity is the important factor in the pathogenesis of gastric ulcer.

METHODS

Gastric acid production was assessed by intermittent aspiration of the stomach after an injection of 0.3 mg. ('standard' test) or 0.6 mg. ('high dosage' test) of histamine (Kay, 1953): the term 'histamine test' denotes the standard test unless otherwise stated. The gastric aspirate was titrated to a pH of 3.5, using Töpfer's reagent, and the output of acid was expressed as milliequivalents over a 90-minute period. The methods for serum vitamin B12 assay and the vitamin B12 absorption and excretion (Schilling) test were as described by Wood, Ralston, Unger, and Cowling (1964). Serological analyses, including the test for complement-fixing autoantibodies to stomach anti-

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3The 90-minute gastric acid production was not measured in test meals before 1956: in these, 'hypochlorhydria' means an acid concentration below 10 mEq. per litre in serial specimens of gastric juice.

CASE DETAILS

The details of the cases studied are described briefly below.

CASE 1 A man, aged 62, with recurring pre-antral gastric ulcer and chronic atrophic gastritis for 15 years, in 1947 presented with epigastric pain of 10 years' duration and intermittent vomiting for three years; radiographs showed 'irregularity' of pyloric region and gastroscopy a small pre-antral ulcer which healed in six weeks.

In 1959 abdominal pain and vomiting recurred; gastroscopy showed a pre-antral ulcer; histamine test produced 0.06 mEq. of HCl.

In 1960 an ulcer was not demonstrable. A gastric biopsy showed advanced chronic atrophic gastritis with a heavy infiltrate of plasma cells and lymphocytes in the lamina propria and patchy intestinal metaplasia of the epithelial surface (Fig. 1), and a high-dosage histamine test, achlorhydria.

Symptoms recurred in 1961 and a pre-antral ulcer was...
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again demonstrated by gastroscopy and radiographs. A high-dosage histamine test gave 0-71 mEq. of HCl.

From 1961 to 1963 there was no further ulceration; the serum vitamin B12 level fell from 290 to 180 μg. per ml.; five gastric biopsies at intervals showed advanced active gastritis with lymphoid infiltration and very scanty pepsin-secreting cells.

The patient died in 1963. There was no necropsy but the presumed cause of death, on clinical evidence, was myocardial ischaemia.

CASE 2 A man, aged 61, had persisting atrophic gastritis after healing of acute gastric ulcer. In 1959 he had had 'indigestion' for seven years, haematemesis and melena for one day. Radiographs and gastroscopy showed a gastric ulcer high on the lesser curvature. A histamine test showed achlorhydria and gastric biopsy pronounced glandular atrophy with dense lymphoid infiltration (Figure 2). The ulcer healed in three months; no recurrence.

The HCl output in three histamine tests during 1959-63 ranged from 0-03 to 7-24 mEq.; three gastric biopsies at intervals showed persisting active atrophic gastritis (Fig. 3), and culture yielded Streptococcus faecalis.

CASE 3 A man, aged 37, with persisting atrophic gastritis and recurrent gastric haemorrhages, in 1949 had mild abdominal pain and haematemesis for five days. A radiograph and gastroscopy were negative; histamine test showed 'hypochlorhydria'. Three gastric biopsies were taken between 1949 and 1951, showing atrophic gastritis with a dense lymphoid infiltration. In 1958-59 he had two gastric bleeds on separate occasions with a recurring small shallow gastric ulcer on the lesser curvature demonstrated by gastroscopy; histamine tests gave 0-73 to 2-28 mEq. Three gastric biopsies at intervals showed severe active atrophic gastritis. In 1959, partial gastrectomy showed an ulcer 3 mm. across on the anterior wall of lesser curvature anteriorly and healed ulcer posteriorly; histology was of widespread atrophic gastritis, lymphoid aggregates, glandular disorganization, numerous eosinophils. Between 1960 and 1964 he had no further haematemeses; histamine test gave 0-15 mEq. A gastric biopsy showed persisting atrophic gastritis; serum vitamin B12 level was 170 μg. per ml.

CASE 4 A man, aged 61, with asthma, had recurrent gastric haemorrhage, gastric ulcer, atrophic gastritis, and early post-gastrectomy vitamin B12 deficiency. In 1949 he had had dyspepsia five years and haematemesis five weeks previously. The histamine test showed 'hypochlorhydria' and gastric biopsy patchy atrophic gastritis. In 1954 he had a large melena but no ulcer was demonstrated. Gastric biopsy showed active atrophic gastritis. In 1956 he had another haematemesis. Radiographs and gastro-
scopv showed a gastric ulcer high on lesser curvature; histamine test gave 0.33 mEq. and gastric biopsy showed advanced atrophic gastritis; later, a perforated gastric ulcer was oversewn. In 1958 after a haematemesis Billroth I gastrectomy was performed; there was an ulcer 1 mm. across on the lesser curvature. Microscopically there was widespread active chronic gastritis (Figure 4). In 1962 the patient had macrocytic anaemia, achlorhydria, advanced gastric atrophy; vitamin B₁₂, 80 µg. per ml.; the Schilling test showed grossly impaired B₁₂ absorption, the 48-hour urinary excretion being 4% of the administered dose.

**CASE 5** A woman, aged 48, was addicted to alcohol and phenacetin with atrophic gastritis, haematemesis, and gastric ulcer. In 1960 she had melena for two days and had had epigastric pains for 10 years. A radiograph and gastroscopy showed a large gastric ulcer on the pars media. A histamine test gave 0.52 mEq. of HCl and gastric biopsy showed atrophic gastritis with large hyperplastic lymph follicle; blood urea was 50 to 66 mg. per 100 ml. In 1961 she had a haematemesis and Billroth I partial gastrectomy was performed. There was an ulcer 1.5 cm. across proximal to the antrum and microscopically widespread active atrophic gastritis. During 1962-64 she had achlorhydria and advanced atrophic gastritis, with biopsy culture yielding *E. coli*.

**CASE 6** A woman, aged 54, had rheumatoid arthritis, Sjögren's keratoconjunctivitis sicca, atrophic gastritis, and gastric ulcer. In 1951 she had had dry, sore eyes, mouth, and tongue for eight years and arthritis for 20 years. Her mother and brother had pernicious anaemia. Examination showed advanced rheumatoid arthritis and keratoconjunctivitis sicca. Between 1951 and 1953 a histamine test showed hypochlorhydria and two gastric biopsies mild superficial gastritis. In 1955 she had painful dry eyes and joint pains, and refractory anaemia. Aspirin consumption was high. A histamine test showed achlorhydria and gastric biopsy advanced active atrophic gastritis.

Epigastric pain and gastrointestinal bleeding were recurring in 1955-56, and a radiograph showed a gastric ulcer. Histamine tests showed 1.15 and 0.56 mEq. of HCl and a gastric biopsy atrophic gastritis (Figure 5). In 1957 Billroth I partial gastrectomy was performed. A penetrating gastric ulcer 1 cm. across was seen on lesser curvature posteriorly and microscopically there was diffuse advanced atrophic gastritis.

In 1958 necropsy findings were rheumatoid arthritis, atrophic gastritis, fibrous replacement of lacrimal glands, normal salivary and thyroid glands, heavy lymphoid infiltration around tracheal glands, and pyelonephritis with renal papillary necrosis.

**CASE 7** A woman, aged 60, had Hashimoto's thyroiditis, chronic hepatitis, atrophic gastritis, and gastric ulcer (reported by Kucers, 1961). In 1958 she had congestive cardiac failure, anaemia, and bleeding gastric ulcer. A thyroid biopsy showed Hashimoto's thyroiditis and liver biopsy active chronic hepatitis with dense lymphoid infiltration. From 1960 to 1963 she suffered from iron-deficiency anaemia, hypothyroidism, cardiac failure, and psychosis. A high-dosage histamine test showed achlorhydria. Gastric biopsy was not performed. Three L.E. cell preparations demonstrated 'free-lying material' and numerous 'rosettes' but no classical L.E. cells. In 1963 necropsy findings were Hashimoto's thyroiditis, cirrhosis, healed gastric ulcer in body of stomach, diffuse chronic atrophic gastritis, aggregations of lymphoid cells in renal cortex and in both adrenal glands, and gross thymic atrophy.

**CASE 8** A woman, aged 54, had a massive haematemesis, numerous acute gastric ulcers, and follicular gastritis. In 1962 after 'mild indigestion' for 20 years, she had a
sudden massive haematemesis and melaena. Polya partial gastrectomy was performed when there were numerous small acute gastric ulcers up to 1 cm. across on the anterior wall near the lesser curvature. Microscopically there was diffuse gastritis, numerous lymphoid follicles with germinal centres in the mucosa, and follicular gastritis (Figure 6). In 1964 she had moderate ‘dumping’, slight macrocytic anaemia (vitamin B12 180 μg. per ml.), achlorhydria on the histamine test. Gastric biopsy showed atrophic gastritis with intense superficial infiltration with plasma cells and some polymorphonuclear cells.

CASE 9 A woman, aged 43, had patchy gastritis and massive gastric bleeding possibly arrested by intravenous prednisolone. In August 1964 after complaining of mild dyspepsia for several years she had gastrointestinal bleeding for two days. Gastroscopy showed a possible shallow gastric ulcer near the antrum. On radiography there was no ulcer in the stomach or duodenum. The histamine test gave 0·50 mEq. of HCl; 25 pints of blood were transfused over 12 days; then prednisolone, 40 mg. given intravenously on three occasions over four days was followed by cessation of bleeding.1 Gastric biopsy showed patchy atrophic gastritis with moderate plasma cell infiltration. Autoantibody tests were negative. In November 1964 she had slight melaena. A histamine test gave 1·48 mEq. of HCl and gastric biopsy showed patchy superficial gastritis.

SEROLOGICAL TESTS AND BIOPSY CULTURES Case 7 gave positive results to tests for autoantibodies to gastric antigen (titre 1/32), thyroglobulin (1/1,280), and cell nuclei; case 6 was not tested and the other cases gave essentially negative results.

1Although it is appreciated that prednisolone per se may cause gastric ulceration, it was given in this case for its ‘anti-inflammatory’ effect. We have not as yet had an opportunity to follow up this observation that prednisolone may have caused arrest of massive bleeding associated with gastritis.

From case 1 we cultured Streptococcus viridans and a micrococcus from the biopsy aspirate, from case 2 Streptococcus faecalis, and from case 5 E. coli. Cases 3, 8, and 9 yielded no growth, and cases 4, 6, and 7, were not tested.

COMMENTS

This paper describes nine patients, four men and five women, with chronic or recurrent gastric ulcers, long-standing dyspepsia attributed to chronic gastritis, and often episodes of severe gastric bleeding which necessitated partial gastrectomy in six. All had impairment of acid production in response to histamine stimulation. Serial gastric biopsies showed persisting gastritis characterized by glandular atrophy and disorganization and heavy infiltration of the lamina propria with lymphocytes and plasma cells, occasionally with formation of lymphoid follicles: polymorphonuclear leucocytes were scanty. In one patient, who bled profusely, the stomach showed numerous lymph follicles, as in follicular gastritis (Magnus, 1952).

We considered the possibility that the gastritis in our patients was of the ‘zonal’ type which is localized to the vicinity of, and perhaps secondary to, a gastric ulcer (Magnus, 1952; Joske, Finckh, and Wood, 1955; Card and Sircus, 1958); however serial gastric biopsies showed that the gastritis persisted despite healing of the ulcer, and gastrectomy specimens in six cases showed the gastritis to be widespread. Moreover Magnus (1952) found that gastritis, often severe in degree with much destruction of gland parenchyma, was present in all of 284 partial gastrectomy specimens removed for gastric ulcer, and this gastritis was diffuse in 26·5% of the cases.

It is our concept that active chronic atrophic gastritis represents the state of ‘decreased mucosal resistance’ or ‘vulnerability’ which is believed to predispose to gastric ulceration (Illingworth, 1956; Kirsner, Clayman, and Palmer, 1959). We would not, of course, exclude a contributory rôle of acid-peptic activity since, in all of our cases, there was some degree of gastric acidity, albeit reduced, and high-dosage histamine stimulation tests occasionally yielded moderate amounts of acid. The biopsy features of the present cases would indicate that ulceration occurs in the active ‘inflammatory’ type of chronic gastritis, which may be patchy ab initio, rather than in the relatively acellular type usually associated with pernicious anaemia, as implied by Washburn and Rozendaal (1938).

The causes of gastritis include dietary irritants, malnutrition, alcohol, and drugs such as aspirin, but the basic pathogenesis of the persisting damage of chronic atrophic gastritis is unknown. The histo-
logical appearance of the stomach in our cases was indicative of an immunological response in the wall of the stomach, and we were interested in the cause(s) of this response and their relationship, if any, to the actual cause(s) of the mucosal damage. One factor considered was chronic infection with enteric bacteria, perhaps facilitated by hypochlorhydria, impaired secretion of mucus, or other undetermined predisposing factors; bacteria were isolated from the stomach in our cases 1, 2, and 5. Another factor could have been autoimmune reactivity, since the chronic atrophic gastritis of pernicious anaemia is associated with serological evidence of autoimmunity to gastric parietal cells (Taylor, Roitt, Doniach, Couchman, and Shapland, 1962; Irvine, Davies, Delamore, and Williams, 1962; Mackay, 1964). It is likely that the autoimmune reaction is the cause, rather than merely a result, of this chronic atrophic gastritis, since, in the comparable incidence of post-gastrectomy gastritis, the incidence of parietal cell antibody is low (te Velde, Abels, Anders, Arends, Hoedemaeker, and Nieweg, 1964). In relation to an autoimmune pathogenesis, case 6 had Sjögren’s disease and rheumatoid arthritis, and case 7 had Hashimoto’s disease, chronic hepatitis, and complement-fixing antibodies to stomach mucosal antigen to a titre of 1/32. We were also interested to note that Doniach and Roitt (1964) reported a raised incidence of parietal cell antibodies in gastric ulcer (22%), but not in duodenal ulcer. However, we would emphasize that in four of our cases we obtained no hint of the cause of the immune response in the stomach nor of the associated mucosal damage.

Our findings suggest that there is a limited group of cases where an established gastritis is followed by gastric ulceration. If the gastritis were initially patchy with mucosal breakdown in one of the involved areas, the more normal areas would be adequate to secrete acid and pepsin which would aggravate and cause persistence of the ulcer. Later the gastritis may become diffuse and involve the entire stomach with depleted acid and pepsin secretion, and yet the ulcer may not heal if the basic pathogenic process responsible for the gastritis persists, be it an extrinsic damaging agent or an autoimmune reaction.

The significance of chronic atrophic gastritis as a disease process was emphasized by Wood and Taft (1958) and by Wood et al. (1964). First, of 221 cases with chronic atrophic gastritis studied in our Unit, 40% suffered from dyspepsia attributable entirely or in part to the gastritis (Wood and Taft, 1958), and chronic gastritis was present in 19% of 200 patients with dyspepsia reported by Coghill (1960): however, some authors are not convinced that gastritis is a cause of symptoms (Coghill, 1960). Secondly, atrophic gastritis is the basic histological lesion of adult pernicious anaemia, but it is uncertain whether ‘simple’ atrophic gastritis inevitably progresses to the gastritis of pernicious anaemia (Williams, Coghill, and Edwards, 1958; Whiteside, Mollin, Coghill, Wynn Williams, and Anderson, 1964): perhaps an additional factor is necessary, i.e., a genetically determined predisposition to form autoantibodies to intrinsic factor or parietal cell antigen. Thirdly, chronic gastritis may predispose to the development of gastric carcinoma, although this is unproven. Fourthly, chronic gastritis, as suggested by our findings and those of others (Langman, Hansky, Drury, and Avery Jones, 1964), may be an important determinant of gastric ulceration and bleeding.

**SUMMARY**

In nine patients a persisting active chronic atrophic gastritis was associated with recurrent or chronic gastric ulceration, often with bleeding. The gastritis was widespread, and persisted or progressed despite healing of the ulcer. Histologically the gastric mucosa showed glandular atrophy, heavy lymphoocyte-plasma cell infiltration, and, frequently, lymphoid follicles, indicative of a sustained immune reaction. These appearances may have been related to bacterial infection or autoimmunity in five of our nine cases: no clear hint as to causation was obtained in the remaining four. We suggest that chronic atrophic gastritis may be an important determinant of gastric ulceration and bleeding.

The vitamin B₁₂ estimations were performed by Dr. D. C. Cowling and Dr. Berta Ugar of the Clinical Pathology Department, Royal Melbourne Hospital. Mr. E. Matthaei, of the University of Melbourne, prepared the photomicrographs. Case 7 was studied through the courtesy of Dr. J. L. Frew.

**REFERENCES**


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