Hypogammaglobulinemia in a family with Crohn’s disease

ROBERT ENNS MD, GUIDO VAN ROSENDAAL MD

Serum immunoglobulins in most Crohn’s disease patients are normal; however, there are occasional reports of Crohn’s patients with hypogammaglobulinemia (1). Low immunoglobulin levels are usually transient, and after appropriate therapy for the disease they increase. We report a case of hypogammaglobulinemia associated with Crohn’s disease. Furthermore, when her family was screened, there was a strong correlation between family members with Crohn’s disease and hypogammaglobulinemia.

CASE PRESENTATION

A 55-year-old female experienced an exacerbation of Crohn’s disease, which was refractory to steroids. Ten years previously the diagnosis was made when recurrent episodes of colicky abdominal pain associated with nausea and vomiting necessitated hospital admission. Barium follow-through examination demonstrated narrowing of the terminal ileum, and she was put on a tapering steroid regimen. Over the next 10 years, biannual exacerbations in her disease were managed with short courses of prednisone and 5-aminosalicylic acid (5-ASA).

Over the year before the reported admission, the patient’s symptoms became increasingly severe and she required a maintenance dose of prednisone 20 mg/day. Despite the addition of metronidazole, nausea, vomiting and right lower quadrant pain persisted, resulting in a 2 to 3 kg weight loss. She had no extra-intestinal manifestations of Crohn’s disease. Medical history included appendectomy, hysterectomy and rheumatic fever. Physical examination was normal except for right lower quadrant tenderness. Family history was significant in that she had six first-degree relatives with Crohn’s disease.

Repeat barium study demonstrated a narrowed segment in
Crohn’s disease and hypogammaglobulinemia

A diagnosis of hypogammaglobulinemia was made, and family screening was done (Figure 1). Four of the six family members with Crohn’s disease had low serum immunoglobulins. Of these four, three had low IgG and one had low IgM. Only two family members without Crohn’s disease were available to have serum immunoglobulins measured and these results were normal. The results demonstrate that the case outlined above was the index case in a family who appear to have coexistent Crohn’s disease and hypogammaglobulinemia (Table 1).

**DISCUSSION**

Gastrointestinal disturbances such as Crohn’s disease have been known to occur in patients with immune-deficient states (2). Some researchers have preferred to call these lesions ‘regional enteropathy-like’ states to denote their similarity to Crohn’s disease (1). Others have simply called the lesions Crohn’s disease and felt that it was associated with hypogammaglobulinemia (3). Despite normal levels of B lymphocytes, patients with hypogammaglobulinemia still have low immunoglobulin levels (4). One explanation is that there is intrinsic abnormality in the B lymphocyte function, which results in low immunoglobulin production. A second theory argues that there is stimulatory dysfunction, so B lymphocytes are not encouraged to produce appropriate levels of immunoglobulins; this theory is supported by several studies that demonstrated that when B cells from affected individuals are isolated and stimulated with specific cytokines, they produce appropriate levels of immunoglobulins. When T cells from affected patients are added to the matrix, immunoglobulin production decreases. Therefore, it appears that the defect in hypogammaglobulinemia may rest in abnormal T cell suppressor activity (4-9).

Only occasional individual reports of hypogammaglobulinemia associated with Crohn’s disease have been published (5,8). This case is unique in that our index patient had six first-degree relatives with Crohn’s disease, four of whom had low levels of immunoglobulins. Three of these four had low serum IgG, and one had low serum IgM and borderline low IgA. These findings are similar to common variable hypogammaglobulinemia, in which it is common to have low IgM, IgA and IgG, especially in siblings. Furthermore, our index case developed chronic obstructive lung disease, common in patients with hypogammaglobulinemia. In some reported cases of hypogammaglobulinemia in Crohn’s disease, serum immunoglobulins returned to normal when the patient had the active disease treated. This did not occur in our index case; she had persistently low serum immunoglobulins after surgical correction and appropriate medical therapy.

**CONCLUSIONS**

The presented index case led us to evaluate further her family, and four other members were subsequently shown to have Crohn’s disease and hypogammaglobulinemia. Serum immunoglobulins measured in the only two available family members without Crohn’s disease were normal. It appears that in this family, Crohn’s disease and hypogammaglobulinemia are closely linked. This finding deserves further study because its elucidation may help to define the pathogenesis of Crohn’s disease more clearly.

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**REFERENCES**
