Familial idiopathic granulomatosis: sarcoidosis and Crohn’s disease in two Indian families

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Abstract
The coexistence of sarcoidosis and Crohn’s disease in different members of the same family is rare and only two instances are on record. Two Indian families showing this association have been studied. In one a brother and sister are affected, and in the other seven and possibly eight persons in two generations have been affected. The familial occurrence of both these conditions supports the view that a transmissible agent may be concerned in the genesis of both diseases in genetically susceptible individuals.

The familial occurrence of a disease may contribute towards a clearer understanding of its pathogenesis, particularly when the aetiology is uncertain. Sarcoidosis and Crohn’s disease are good examples. These conditions have been considered to be readily distinguishable from each other1 but recent research suggests that this may not be so.2 Some of the features that were thought to occur in only one condition have now been observed in both.1,3 Sarcoidosis and Crohn’s disease1,2 are rare in India but are now being seen with increasing frequency.3-4 They may have been mistaken for tuberculosis in the past.3 The coexistence of these two uncommon conditions in the same family has been recorded only twice.5-6 We describe two Indian families where sarcoidosis and Crohn’s disease have affected different members.

Case reports
FAMILY 1
Patient 1 A 50 year old housewife developed a low grade intermittent pyrexia (37.5–38.5°C), anorexia, and lethargy in 1980. Physical examination and laboratory tests showed nothing abnormal but the fever persisted. In 1981 she had a documented lower urinary tract infection, which responded to antibiotics. In 1982 she developed erythema nodosum, arthralgia, and severe anterior and posterior uveitis, which responded to corticosteroids but flared up whenever an attempt was made to reduce the dose. Bilateral hilar lymphadenopathy was noted on her chest radiograph and later the same year a cervical gland biopsy showed discrete non-caseating epithelioid cell granulomas with occasional giant cells. As the histological features resembled tuberculosis she was treated with isoniazid, pyrazinamide, and ethambutol for 18 months even though the tissue was negative for acid fast bacilli on stains and in culture. While having this treatment she developed bilateral parotid swelling and a left seventh cranial nerve palsy. A diagnosis of sarcoidosis was considered and a Kveim test gave a positive result. Disease activity continued until 1985 and then subsided. She is currently symptom free and has not received corticosteroids for five years.

Patient 2 The 56 year old brother of patient 1 developed fever, abdominal pain, and an altered bowel habit in 1978. Initial examination showed nothing remarkable but a discrete, firm lump became palpable in the caecal region in 1979. At colonoscopy the caecum was deformed, inflamed, and ulcerated but there was no evidence of malignancy in several biopsy specimens. Suspected of having amoebiasis, he was treated with metronidazole and had some relief. He had severe pain, colic, distension, vomiting, and right lower quadrant tenderness, however, in 1980, and underwent an emergency laparotomy elsewhere. The terminal ileum was found to be thickened, rigid, indurated, and encased in mesenteric fat. The paracolic region had several pockets of pus, which were drained. No resection was carried out. He was advised to take isoniazid, ethambutol, and thiacetazone for 18 months. He remained well for a few months but symptoms recurred in 1981, when he had several episodes of subacute intestinal obstruction with fever. Repeat colonoscopy showed aphthous ulcers in the caecum and several biopsy specimens showed granulomatous colitis consistent with Crohn’s disease. No further treatment was prescribed. Since then he has been well except for mild colic on several occasions.

FAMILY 2
This is a large family with eight affected members in two generations (figure). There has been no consanguinity and the origin of the family is entirely north Indian. Only one member of the family at present resides in India (patient 8) and the details of her illness are given below. As the others have not been examined by us personally their clinical problems are summarised.

Patient 1 Died aged 45 years from a chronic respiratory illness and cor pulmonale. No histological specimens were obtained.

Patient 2 Died aged 60 years of an illness very similar to that of his brother (patient 1). In addition he had dermal plaques and nodules, from which a tissue diagnosis of

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sarcoidosis was made before he died.

*Patient 3* is living and being treated for dermal, pulmonary, and cardiac sarcoidosis in the United States. The diagnosis has been confirmed histologically. She is 72 years old. 

*Patient 4* Died aged 70 years. She had had histologically confirmed sarcoidosis of the skin, eyes, and lungs for several years. 

*Patient 5* is aged 43 and resident in England. He has a permanent ileostomy for severe Crohn’s disease. 

*Patient 6* is the 40 year old brother of patient 5, and is also resident in England. He also has an ileostomy for Crohn’s disease. In addition, he had nodular skin lesions on the legs.

*Patient 7* is 44 years old and living in Canada. He has had ill health with fever, lymphadenopathy, and cough, but does not have histologically confirmed sarcoidosis. The tuberculin test response was negative. 

*Patient 8*, a 40 year old housewife, developed thickening, redness, and dryness of the skin over the malar region in 1986. The lesion was mildly photosensitive. She also had cervical lymphadenopathy. Skin biopsy showed discrete epithelioid cell granulomas with occasional giant cells, no caseation, and no acid fast bacilli. Her chest radiograph and spirometric indices were normal, but she had hypercalcaemia and hypercalciuria. She responded well to corticosteroids and attempts to taper the dose for the first three years were followed by a relapse. For the last year the disease appears to be quiescent, though in 1989 she passed a urinary calculus. At present she is taking a small, alternate day dose of prednisolone.

**Discussion**

Although familial sarcoidosis is well described the mode of inheritance is not clear.17-34 HLA studies in different ethnic populations have shown variable associations.16 The B8/CW7/DR3 haplotype has been found to be related to acute articular, dermal, and ocular sarcoidosis, with a good prognosis in English patients.11 Familial aggregates of A2/B7/CW7/DR2/DR7 and A2/B5/CW7/DR5 have been reported.12 13 Family studies in inflammatory bowel disease have clearly established that a familial incidence is far greater than would be expected by chance.14 

Sarcoidosis and Crohn’s disease were believed to be easily distinguishable granulomatous conditions of unknown aetiology. Whereas the former is usually widely disseminated, the latter is considered to be limited to the bowel predominantly. Bowel lesions in sarcoidosis are very infrequent even at necropsy.1 Evidence, however, suggests that the two conditions may have more features in common than has been recognised, despite the fact that the two pioneers, Crohn and Siltzbach, were both working at Mt Sinai Hospital in New York. The extraintestinal manifestations of Crohn’s disease show some similarity to some of the dermal, articular, and ocular presentations of sarcoidosis. Lymphocytic alveolitis, long considered a hallmark of sarcoidosis, has been shown to occur in Crohn’s disease.2 There have been well documented reports on patients with both diseases15-17 and on at least two families18 in which the two conditions have coexisted.

The interplay of the genetic makeup of the individual with the environment in the development of disease has been strongly supported by several studies. Acute sarcoidosis has been documented in close relatives who have lived together, and this has been used as evidence for the existence of a transmissible agent responsible for the disease.18 There has been a great resurgence of interest in the idea that mycobacteria may be responsible for both sarcoidosis19-20 and Crohn’s disease.21 22 This has been based on the detection of proteplastic forms of mycobacteria in tissue in both conditions.20 22 Possibly the two conditions are manifestations of the same disease, which occurs in susceptible persons and is modulated by a genetically determined abnormal immunological response to mycobacteria. The seemingly different clinical features could be explained by different routes of entry of the antigenic stimulus—inhalation for sarcoidosis
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and ingestion for Crohn’s disease. The reported familial clustering of these conditions supports this contention.

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