Behçet's syndrome: a family study and the elucidation of a genetic role

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SUMMARY

A family with features of the mucocutaneous ocular syndrome is described. A 4-generation study reveals that the condition has been inherited as an autosomal dominant trait with variable expressivity. Psychiatric symptoms and myalgia have been prominent features peculiar to our patients.

The aetiology of Behçet's syndrome has been disputed ever since Behçet's description (Behçet, 1937; Behçet, 1940). Both viral (Behçet, 1937; Sezer, 1952, 1953, 1956; Evans et al., 1957; Dudgeon, 1961) and autoimmune (Jensen, 1941; Shimizu et al., 1965) mechanisms have been postulated and familial occurrence has been reported (Fadli and Youssef, 1973; Fowler et al., 1968; Mason and Barnes, 1969; Sezer, 1956). Goolamali et al. (1976) presented a 4-generation family study of Behçet's syndrome associated with psychiatric disorders, and a genetic transmission was postulated.

Patients

We present a 4-generation family study, 15 of whose members fulfil Curth's (1946a, b) criteria of Behçet's syndrome. Five additional members, all under 10 years of age, exhibit mild expression of the symptom complex. Some of the patients presented with the more unusual manifestations of the syndrome, myalgia and psychiatric symptoms.

A pedigree of the family is presented in Fig. 1, from which it can be seen that of 42 family members personally examined, 15 fulfilled Curth's criteria for the diagnosis of Behçet's syndrome, namely 2 of the 3 principal lesions of the triple symptom complex: aphthous stomatitis, genital ulceration and ocular inflammation. Ten of the 15 affected individuals are female and the age at onset of the disease varied between 2½ years and 50 years. Five additional members (3 males, 2 females), all aged 10 years or younger, have oral ulceration only, which is probably the earliest manifestation of the symptom complex. The affected mother of the index patient (I-5) was of Huguenot descent. There was no consanguinity and no history of any obvious

Fig. 1. Pedigree of the family.
beh"et's syndrome: a family study and the elucidation of a genetic role

Table 1 Clinical findings encountered among 20 affected family members

<table>
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<th>Patient</th>
<th>Sex</th>
<th>Age</th>
<th>Clinical features</th>
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*Pericarditis. **Bilowing posterior mitral leaflet. †Died.

environmental agents operating to produce the clinical findings in the family.

Oral Lesions

Painful recurrent oral ulcers were recorded in all patients (Table 1). These occurred singly or in crops on the oral mucosa, lips, tongue, pharynx, soft palate, larynx, and in the nasal mucosa. They persisted for days to months, and their size varied from pinhead to 2 cm diameter. Emotional stress was implicated as a precipitating factor in 8 patients.

Genital Ulcers

Eight females and 4 males had painful single or multiple genital ulcers which resembled the oral ulcers both in appearance and in course.

Ocular Lesions

Four females and 3 males had ocular problems. All had evidence of iridocyclitis. Corneal ulcers were reported in 4. Patient III–3 is unusual in being affected at the early age of 7 years.

Skin Lesions

The index patient (II–9), 42 years, has severe acne and numerous generalised skin postules. Two children aged 2 ½ years (VI–1) and 4 years (IV–6) have eczema.

Myalgia and Arthralgia

Myalgia and arthralgia were prominent in 9 patients. The pain was described as deep-seated and gnawing in all muscles, but more particularly in the arms and chest wall. Movement and pressure did not aggravate the discomfort, and rest did not ease the pain. Simple analgesics and anti-inflammatory agents were ineffective. The pain could last for weeks. The severity and duration of the pain induced the index patient (II–9) to become dependent on habit-forming drugs. Colchicine, however, was discovered to produce a dramatic improvement of the myalgia within 24 hours of taking the drug. Relapse would occur immediately after stopping the drug. The other symptoms were marginally improved by Colchicine. Arthritis was not a prominent manifestation in any of the patients.

Psychological disturbances

Ten patients had mental depression requiring therapy. The index patient (II–9) was committed to a mental institution. A woman aged 24 years (II–7) committed suicide.

Cardiovascular system

A man (I–I) aged 43 years had an unexplained pericarditis, and patient IV–4, a boy of 5 years, had recurrent attacks of paroxysmal atrial tachycardia. Patients II–6 and II–9 have Barlow’s syndrome (billowing posterior mitral leaflets).

Discussion

Arthritis, although well described in Behçet’s syndrome (Mason and Barnes, 1969) was not present in our patients. However, 9 patients presented with severe muscle pain, incapacitating them and lasting for intervals of weeks before remission. This symptom has not previously been described. Colchicine
was found to have a dramatic effect on the alleviation of the pain. Since the basic pathological lesion is considered to be a vasculitis of small vessels (Nazzaro 1966; O’Duffy and Carney, 1971; Shikano, 1966, Lehner, 1969; Saito et al., 1971), with a perivascular lymphmononuclear cell infiltration and an exaggerated chemotactic polymorphonuclear response (Mizushima et al., 1977), it is possible that this may be explained by its inhibition of polymorph chemotaxis (Mizushima et al., 1977).

Psychological symptoms of depression requiring therapy were found in 10 patients. One patient committed suicide. Neurological signs were absent. Dementia has been described in patients with neurological signs, but there have been few reports of psychiatric manifestations of Behçet’s syndrome. Schotland et al. (1963) referred to 9 patients in whom signs of confusion and emotional lability were present. In a psychiatric study of Behçet’s syndrome (Epstein et al., 1970) submissiveness, depression, and neuroses were observed. Goolamali et al. (1976) described the development of acute schizophrenia with affective features in one of their patients.

Two patients were found to have Barlow’s syndrome. The unaffected members were examined but not found to have this entity. As this condition is relatively common, it is likely to be independent of Behçet’s syndrome.

Chajek and Fainaru (1975) reviewed Behçet’s syndrome and emphasised the male preponderance. Ten of our patients were female. Our youngest patient (III-3) was 7 years and our oldest died at the age of 70 years (I-5). Five children (Table 1), all younger than 10 years, have aphthous ulcers.

Published reports of genetic transmission of Behçet’s syndrome are rare. Sezer (1956) diagnosed the syndrome in 3 brothers and Fowler et al. (1968) described neurological manifestations in 2 sisters with Behçet’s syndrome.

Mason and Barnes (1969) reported 28 cases, and 4 of these had affected family members: a man had 1 brother who was definitely affected and another who was possibly affected; a woman’s father and 2 siblings had buccal ulcers (while her mother possibly had Behçet’s syndrome); another woman’s mother was possibly affected; and an affected woman had 1 daughter definitely and another possibly affected. In the discussion which followed the presentation of Mason and Barnes’s paper, D. A. Pitkeathly of Wigan, described a family in which brother and sister had Behçet’s syndrome, while their mother had atypical arthritis with iritis, ‘But one couldn’t make the diagnosis of Behçet’s syndrome definitely; she, however, has unilateral sacroillitis’. Chajek and Fainaru (1975) encountered among their 34 predominantly Jewish patients 3 pairs of brothers with Behçet’s syndrome; 1 pair had an affected sister and a nephew with recurrent aphthous stomatitis. Chajek et al. (1977) again referred to these 2 families and in addition reported the significant association between Behçet’s syndrome and HLA B5. Fadl and Youssef (1973) reviewed 46 cases with Behçet’s syndrome in the United Arab Republic and showed that in 3 instances, 2 siblings had the disorder. Goolamali et al. (1976) have reported a family in which 1 individual has been involved in each of 3 generations, and 2 individuals in 1 generation, compatible with autosomal dominant inheritance. The HLA haplotype 1–17 was present in all 4 affected persons.

It appears that the condition we have described is inherited as an autosomal dominant condition with variable expressivity, even though there is no male to male transmission. There is 1 patient (II-5) (Fig. 1) in whom the gene would appear to be non-penetrant.

If one postulates that Behçet’s syndrome is inherited as an autosomal dominant with variable expressivity (and even occasional non-penetrance), then the few previous family reports would lend support to this theory. There may, in fact, be nothing unique about the family reported here, and if this is so it may be expected that, as clinicians specifically search for affected relatives, more families will be reported.

References


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