Behcet’s syndrome in 32 patients in Yorkshire

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SUMMARY Thirty-two patients with Behcet’s syndrome were surveyed clinically, radiologically, and immunologically. HLA antigens were determined in a proportion. The patients were culled from a defined geographical area (Yorkshire) with a population of 5 million. 21 patients satisfied Mason and Barnes’s (1969) criteria for major disease. 100% had mouth ulceration, 91% genital ulceration, 66% skin complaints, and 63% arthritis (of the nonerosive type previously documented). 3 patients had gastrointestinal involvement, one of whom had proven Crohn’s disease. One patient had psoriasis. One patient had a popliteal artery aneurysm resected and one has since had surgery after the development of a tracheo-oesophageal fistula. No sacroilitis was found.

Behcet’s syndrome is an uncommon disorder first described by Professor Behcet in 1937 but with a long pedigree possibly going back to Hippocrates. There is a substantial ophthalmological and dermatological literature dating from the turn of the century, the two streams converging over the past 20 years. Information has accumulated at an increasing rate over the last few years and has included several large series and reviews (Chajek and Fainaru, 1975; Zizic and Stevens 1975). Classically Behcet’s syndrome is considered to consist of a triad of findings (oral and genital ulceration and iritis) but many authors have widened the scope of the diagnosis and varied the diagnostic criteria. Curth (1946) suggested that the diagnosis could be made when two features of the triad were present. The many names for different groups of features found in Behcet’s syndrome have led to confusion in diagnosis. Dowling (1961) commented on the difficulties of establishing a certain diagnosis. Recently in an attempt to clarify the position, Mason and Barnes (1969) suggested a series of major and minor criteria.

Originally the disease appeared largely confined to the Middle East but recently reports have come from all parts of the world, particularly from Japan. One of the few epidemiological reports suggests that the disease occurs in 1 in 10 000 of the population of Hokkaido, a remote part of Japan (Aoki et al., 1971).

Work done in the unit at Leeds over the past 10 years has led to the concept of the seronegative spondarthropathies. Behcet’s syndrome has been tentatively placed within this orbit as there are certain features of the disorder which are also found in ulcerative colitis and ankylosing spondylitis (for example, erythema nodosum, mouth ulceration, and iridocyclitis). It may be that these features all arise on the basis of a common genetic background. Studies of histocompatibility (HLA) antigens offer one means of looking at this in more detail. The incidence of HLA B27 is raised in ankylosing spondylitis, as it is in patients with uveitis from other causes.

The work reported here is part of a family study of Behcet’s syndrome in the county of Yorkshire. Analysis of probands aimed to define the major components of the disorder as presenting in a defined geographical area and to determine the frequency of the disease in that area, as well as the prevalence of sacroilitis and the various HLA antigens. The presence of coincident diseases associated with seronegative arthropathies has been noted in an attempt to learn more about the nature of Behcet’s syndrome and its relation to the seronegative arthropathies.

Patients

SOURCE

All general practitioners in Yorkshire were contacted via the appropriate executive councils for the names of patients with a possible diagnosis of Behcet’s syndrome and for permission to approach them. Hospital colleagues in the region were also contacted. A standard letter was sent to each of those named inviting them to participate in the survey.
Those willing were questioned and examined if they satisfied the following criteria. (1) Currently resident in the County of Yorkshire. (2) Mouth ulceration had been recurrent over at least a 6-month period. (3) Another feature of the disease was present.

**EXAMINATION**

The method used was substantially that outlined by Moll (1971) with a questionnaire adapted for Behçet's syndrome. A standard proforma was designed to give details of the various manifestations of the disease. Thus, skin complaints such as psoriasis, acne, erythema nodosum, and infective lesions were sought. Detailed features of mouth and genital ulceration were noted and patients with Reiter's disease excluded. Conjunctivitis, iritis, backache, sacroiliitis, arthritis, thrombosis, and neurological abnormalities were all recorded.

A full clinical examination was supplemented by measurement of chest expansion and lumbar spinal movement, as described by Moll et al. (1972) and Moll and Wright (1972). Blood was taken for an immunological profile, haemoglobin, erythrocyte sedimentation rate, and white cell count. HLA antigen profiles were also undertaken on a substantial number of patients by Dr. S. M. Rajah of the Leeds Blood Transfusion Unit. Radiographs of the sacroiliac joints were obtained from males over the age of 15 years and females over the age of 45 years, according to MRC criteria and using a 4" gonad protector. The hands, feet, and any affected joints were x-rayed.

**Methods**

**IMMUNOFLOURESCENCE STUDIES**

**Detection of autoantibodies**

Antigens were provided by unfixed cryostat sections of human kidney, stomach, and thyroid, mounted as one block, and by sections of guinea pig lip and oesophagus mounted as a second block. Sera were reacted with human tissues at an initial dilution of 1:4 and with guinea pig tissues at an initial dilution of 1:15, using standard indirect immunofluorescent tests. For these tests multivalent antihuman immunoglobulin antisera labelled with fluorescein isothiocyanate were used at dilutions of 1:64. Sections were examined under incident illumination on a Leitz Orthoplan microscope, using narrow band excitation (495 mm) and a K530 barrier filter.

**Immunoglobulins**

IgG, IgM, and IgA were analysed quantitatively and the values obtained recorded in a graded fashion for computer analysis as follows:

<table>
<thead>
<tr>
<th></th>
<th>IgG (g/l)</th>
<th>IgM (g/l)</th>
<th>IgA (g/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 unknown</td>
<td>&lt; 9·8</td>
<td>&lt;0·51</td>
<td>&lt;1·67</td>
</tr>
<tr>
<td>1 below normal</td>
<td>9·8-15·2</td>
<td>0·51-1·09</td>
<td>1·67-3·09</td>
</tr>
<tr>
<td>2 normal range</td>
<td>15·2-20·1</td>
<td>1·10-2·0</td>
<td>3·10-3·50</td>
</tr>
<tr>
<td>3 slightly raised</td>
<td>&gt;20·1</td>
<td>&gt;2·0</td>
<td>&gt;3·50</td>
</tr>
<tr>
<td>4 greatly raised</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**PREPARATION AND ANALYSIS OF DATA**

Data were analysed conventionally by calculating the percentage incidence of various characteristics. Data were also prepared for Q-analysis (Atkin 1974) as described fully elsewhere (Chamberlain, 1976). Q-analysis provides a method of studying the connectivities between members of a set Y when these are related (via a mathematical relation) to another set X, in this case: Y = (names of individual people), X = (names of disease attributes).

Q-analysis is a dynamic and flexible process, one of its merits being that no initial assumptions have to be made as to which features in a large mass of data are associated individually or in subgroups. A consistent coding principle was adopted for the computer analysis: absence of data was indicated by 0, normal values were presented by unity, and increasing pathology indicated by progressively higher digits. This allowed study of the connection between disease attributes at increasing levels of diagnostic certainty.

In addition the patients could also be stratified in several established ways in this survey. 3 patients showed the classical triad of features. Using Mason and Barnes's (1969) criteria (but excluding the family history, as the value of this was under investigation; and cardiovascular disease, as no details of this were available), 21 patients satisfied the criteria for definite disease. Radiographs of the sacroiliac joints were graded as described by Macrae et al. (1971).

**Results**

Forty-one subjects were contacted, 33 agreed to take part in the survey; 1 of these was rejected as not satisfying the entry criteria and thus the number with Behçet's disease entering the survey was 32 (Table 1). (17 were referred from general practitioners and 15 from hospitals.) The group was composed of 20 females and 12 males in whom the mean age of onset was 24-7 years, only 4 patients developing the disorder after the age of 40 years (Table 2). Mouth ulceration was present in all cases, by definition, being the presenting feature in 25 patients; in 16 it preceded all other symptoms. Arthritis was the earliest feature in 3, skin complaints in 10, and genital ulceration in 2 (Table 3). The major clinical features of the disorder are shown in Table 4, in
Table 1 32 patients diagnosed as having Behcet's syndrome

<table>
<thead>
<tr>
<th>Case no.</th>
<th>M/F</th>
<th>MU</th>
<th>GU</th>
<th>Iritis</th>
<th>Skin lesions*</th>
<th>GI</th>
<th>Vascular†</th>
<th>Arthritis‡</th>
<th>CNS</th>
<th>SCAT</th>
<th>HLA B27</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>30</td>
<td>M</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>A, P</td>
<td>T</td>
<td>+</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td></td>
<td>R macular haemorrhage</td>
</tr>
<tr>
<td>31</td>
<td>M</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>A, S, E, M</td>
<td></td>
<td></td>
<td>+</td>
<td>0</td>
<td>0</td>
<td></td>
<td>R popliteal artery aneurysm</td>
</tr>
<tr>
<td>32</td>
<td>M</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>E</td>
<td>Ar</td>
<td>+</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Patients with definite Behcet's syndrome (Mason & Barnes, 1969)

| 1 | F | + | + | C | A | T | + | 0 | 0 | Rectal and anal ulcers |
| 2 | F | + | + | C | A, S, E, M | + | 0 | 0 | Arytenoid ulcer |
| 3 | M | + | + | C | A, S, E, M | + | 0 | 0 |                      |
| 5 | F | + | + | P |   |   | 0 | NT |   |                      |
| 6 | M | + | + | A, S | + | 0 | 0 | NT |   |                      |
| 8 | F | + | + | E |   |   | 0 | 0 |   |                      |
| 9 | M | + | + | A, S, E, P | T | 0 | 0 |   |   |                      |
| 10 | M | + | + | S | T | 0 | 0 | NT | Tracheo-oesophageal fistula |
| 12 | M | + | + | A, S, Ps, E | + | 0 | NT | Psoriasis, died |
| 14 | F | + | + | A, S | T | 0 | 0 |   |                      |
| 15 | M | + | + | S, E | T | 0 | 0 | NT | Round cell infiltrate in thickened vascular endoepithelium of arterioles |

Patients with possible Behcet's syndrome

| 4 | F | + | + | C | OA | 0 | 0 |   | Died quadriplegic |
| 7 | F | + | + |   | OA | 0 | 0 |   |                      |
| 11 | F | + | + |   | + | 0 | 0 |   |                      |
| 16 | F | + | + |   |   | 0 | NT |   |                      |
| 18 | F | + | + |   |   | 0 | 0 |   |                      |
| 19 | M | + | 0 | C | A, S | 0 | 0 | Aged 11 |                      |
| 20 | F | + | + |   |   | 0 | 0 | Aged 15 |                      |
| 21 | F | + | + |   |   | 0 | 0 |   |                      |
| 24 | F | + | 0 | T |   | 0 | 0 | Recurrent vaginitis |                      |
| 27 | F | + | + |   | RA | 0 | 0 |   |                      |
| 33 | M | + | 0 | + |   |   | 0 |   |                      |

Total 32 29 4 21 3 8 20 8 1 6

MU, GU = mouth and genital ulcers; C = conjunctivitis (recurrent); GI = gastrointestinal; SCAT = sheep cell agglutination test; + = positive; NT = not tested.

* = acne; S = sepsis; Ps = psoriasis; E = erythema nodosum; M = maculopapular rash; P = sepsis on pin prick (recurrent); U = ulcers on skin.

†T = thrombophlebitis; Ar = arterial lesions.

‡A = destructive arthritis; OA = arthritis with x-ray evidence of osteoarthrosis; RA = clinical rheumatoid arthritis.

Table 2 32 patients diagnosed as having Behcet's syndrome in Yorkshire: onset of disease by decade

<table>
<thead>
<tr>
<th>Years</th>
<th>No. of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-10</td>
<td>8</td>
</tr>
<tr>
<td>11-20</td>
<td>8</td>
</tr>
<tr>
<td>21-30</td>
<td>5</td>
</tr>
<tr>
<td>31-40</td>
<td>7</td>
</tr>
<tr>
<td>41-50</td>
<td>1</td>
</tr>
<tr>
<td>51-60</td>
<td>2</td>
</tr>
<tr>
<td>61+</td>
<td>1</td>
</tr>
</tbody>
</table>

which it will be seen that genital ulceration and arthritis were dominant.

One patient had psoriasis, 1 suffered from Crohn's disease leading to colectomy, 1 complained of perianal ulceration, and a further patient had a caecectomy for nonspecific caecal ulceration. 2 patients died during the survey. One, a patient with late-onset Behcet's disease who was treated for this and for severe psoriasis with high dosages of corticosteroids, died of Gram-negative septicemia. The other had been quadriplegic and dysarthric due to long-standing Behcet's disease and died of uraemia and large necrotic bedsores during the survey.

Table 3 Onset of Behcet's syndrome: presenting features

<table>
<thead>
<tr>
<th>Years</th>
<th>No. of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-10</td>
<td>8</td>
</tr>
<tr>
<td>11-20</td>
<td>8</td>
</tr>
<tr>
<td>21-30</td>
<td>5</td>
</tr>
<tr>
<td>31-40</td>
<td>7</td>
</tr>
<tr>
<td>41-50</td>
<td>1</td>
</tr>
<tr>
<td>51-60</td>
<td>2</td>
</tr>
<tr>
<td>61+</td>
<td>1</td>
</tr>
</tbody>
</table>

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Table 4  Characteristics of patients with Behcet’s syndrome in Yorkshire

<table>
<thead>
<tr>
<th></th>
<th>M (n = 12)</th>
<th>F (n = 20)</th>
<th>Total (n = 32)</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gastrointestinal disorder</td>
<td>0</td>
<td>3</td>
<td>3</td>
<td>9</td>
</tr>
<tr>
<td>Vasculitis</td>
<td>4</td>
<td>4</td>
<td>8</td>
<td>25</td>
</tr>
<tr>
<td>Neurological complaints</td>
<td>3</td>
<td>5</td>
<td>8</td>
<td>25</td>
</tr>
<tr>
<td>Eye complaints (all)</td>
<td>5</td>
<td>3</td>
<td>8</td>
<td>25</td>
</tr>
<tr>
<td>Skin complaints (all)</td>
<td>11</td>
<td>10</td>
<td>21</td>
<td>66</td>
</tr>
<tr>
<td>Arthritis</td>
<td>10</td>
<td>10</td>
<td>20</td>
<td>63</td>
</tr>
<tr>
<td>Genital ulcers</td>
<td>10</td>
<td>19</td>
<td>29</td>
<td>91</td>
</tr>
<tr>
<td>Mouth ulcers</td>
<td>12</td>
<td>20</td>
<td>32</td>
<td>100</td>
</tr>
</tbody>
</table>

MOUTH ULCERATION
Many authors have commented that mouth ulceration in Behcet’s syndrome has no specific features and that the individual ulcers may be aphthous or herpetiform. This is undoubtedly true when the syndrome is mild but when florid the picture is dramatic and even diagnostic, ulcers extending from the lips to the fauces and into the oesophagus where they cause substernal pain and difficulty in swallowing. At this stage ulcers are usually numerous, large, deep, and may even result in scarring. One patient had surgery to relieve subsequent circum-oral contractures. Another was found to have an arytenoid ulcer and one male in our series has recently developed a tracheo-oesophageal fistula, secondary to Behcet’s ulceration. Frequently the ulcers are so painful that the patients only eat liquid foods, after the application of local anaesthetic. Ulcers may be present almost continuously with only short intervals between attacks and with little periodicity in relation to other features of the disorder or to the menses. On the other hand, they are often infrequent, small, and scantly.

Various medications were tried by all patients and indeed were so numerous and varied that no statistical analysis of their effect has been attempted.

GENITAL ULCERATION
Genital ulceration was present in almost all patients (10 male, 19 female) often beginning shortly after mouth ulceration. Either the mucosa or the skin of the area was affected. Vaginal or cervical ulceration was frequently detectable only by presence of a vaginal discharge, bleeding, or dyspareunia, the ulcerated area often being painless although tender on contact; vulval ulceration occasionally led to labial scarring and destruction.

Discussion showed that the possibility of the disease being venereal had often been an unresolved source of anxiety for the proband and/or their spouse. Indeed, 10 patients with genital ulceration had experienced depression, made suicidal bids, or had suffered marriage breakdown.

SKIN ABNORMALITIES AND SEPSIS (Table 5)
It was our impression that sepsis was frequent but usually minor in these patients. Acne was present in 10 (and conjunctivitis and styes were commonly reported). Erythema nodosum had occurred at some stage in the disorder in 8, and other conditions such as maculopapular rashes were noted in 3 (Table 7). One patient had psoriasis. The development of pustular lesions at the site of needle puncture had been commented on by Blobner (1937) and others and were thought to be diagnostic of the condition. A history of such lesions was given by 5 of our patients and was never recorded from spouses or relatives. The relationship of sepsis to the syndrome has been commented on by Behcet himself but has never been clear.

ARTHRITIS (Table 6)
Twenty patients in this series suffered from some form of arthritis (the mean age of onset being some 2 years after the beginning of the disease itself: mean 27.4 years, range 5–64 years). Of 15 patients judged to have Behcet's arthritis, 3 experienced arthralgia and 12 had documented synovitis which was often spasmodic but not truly migratory and left no sequelae, either radiological or clinical.

Table 5  Behcet’s syndrome: skin conditions

<table>
<thead>
<tr>
<th></th>
<th>Total (n = 32)</th>
<th>M (n = 12)</th>
<th>F (n = 20)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acne</td>
<td>10</td>
<td>7</td>
<td>3</td>
</tr>
<tr>
<td>Psoriasis</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Erythema nodosum</td>
<td>8</td>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td>Maculopapular rash</td>
<td>3</td>
<td>1</td>
<td>2</td>
</tr>
</tbody>
</table>

Table 6  15 patients with nonerosive arthritis of Behcet’s syndrome

<table>
<thead>
<tr>
<th>Joint</th>
<th>Presenting joint (no. of patients)</th>
<th>Dominant affected joint (no. of patients)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shoulder</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Elbow</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Wrist</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Metacarpophalangeal</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Proximal inter-phalangeal</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Total upper</td>
<td></td>
<td></td>
</tr>
<tr>
<td>limb joints</td>
<td>8</td>
<td>5</td>
</tr>
<tr>
<td>Hip</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Knee</td>
<td>3</td>
<td>6</td>
</tr>
<tr>
<td>Ankle</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Metatarsophalangeal</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Interphalangeal</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Total lower</td>
<td></td>
<td></td>
</tr>
<tr>
<td>limb joints</td>
<td>7</td>
<td>10</td>
</tr>
</tbody>
</table>
these 15, the shoulder and knee joints were first.
affected and large joints were affected more
commonly than small joints (Table 6). 2 patients suffered
a nonarthritis of the knee. None had ankylosing
spondylitis or gout. The patient with psoriasis
(Case 12) had radiological evidence only of osteo-
arthrosis of the toes and clinically a palindromic
polyarthritis of the knees, elbows, and toes which
was thought to be due to Behcet's syndrome, since it
was coincident with other features of Behcet's
disease.

Of the 5 further patients, 3 had clinical rheumatoid
arthritis substantiated in 2 cases by nodules and
subluxed joints and in 1 by seropositivity (x-rays
unavailable), and the remaining 2 cases had clinical
and radiological evidence of osteoarthritis.

BACKACHE (Table 7)

Eleven patients complained of lumbar pain at some
stage, 3 also experiencing pain in thoracic or
cervical regions. The mean age of onset of pain was
35 years (range 19–35 years). In all but 2 pain had
been short-lived (2 weeks or less) and appeared to
contribute little to the suffering experienced by
patients in the course of Behcet's disorder, even
though sciatica accompanied lumbar pain in 60.
2 of the 3 patients with low back pain and reduced
lumbar movement, according to Moll's criteria, had
normal radiographs of the SI joints. The third was
not x-rayed.

EYE INVOLVEMENT

Recurrence conjunctivitis was found in 4 patients and
was of little importance. In contrast, where irido-
cyclitis had occurred (4 males) it was severe, chronic,
and a prominent feature of the disease which was
difficult to bring under control.

NEUROLOGY

While a quarter of the patients admitted on direct
questioning to having had paraesthesiae and numbness, no abnormalities were detected clinically
at the time of interview except in one patient
(Case 29) who had significant neurological involve-
ment and died at the age of 40 years of uremia
(see above)

VASCULAR INVOLVEMENT

Eight patients (4 male, 4 female) had vascular
abnormalities. 7 had experienced thrombophlebitis

Table 7  Behcet's syndrome: chest expansion and spinal movements in probands

<table>
<thead>
<tr>
<th>Case no.</th>
<th>M/F</th>
<th>Chest expansion</th>
<th>Lumbar spinal movement</th>
<th>Pain site</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>&lt; 2.5 cm</td>
<td>Reduced*</td>
<td>Forward flexion</td>
</tr>
<tr>
<td>3</td>
<td>M</td>
<td>–</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>6</td>
<td>M</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>7</td>
<td>F</td>
<td>+</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>22</td>
<td>F</td>
<td>–</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>23</td>
<td>F</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>27</td>
<td>F</td>
<td>–</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>32</td>
<td>M</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Total</td>
<td>4F:3M</td>
<td>2</td>
<td>2</td>
<td>4</td>
</tr>
</tbody>
</table>

*J. Moll’s criteria.

Table 8  Phlebitis (7) and arteritis (1) in Behcet's syndrome (32 patients): characteristics of patients with vascular involvement

<table>
<thead>
<tr>
<th>Case no.</th>
<th>M/F</th>
<th>MU</th>
<th>GU</th>
<th>Conjunctivitis</th>
<th>Iritis</th>
<th>Skin</th>
<th>Erythema nodosum</th>
<th>Arthritis</th>
<th>Neurology</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>F</td>
<td>+</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>Acne</td>
<td>+</td>
<td>0</td>
<td>+</td>
</tr>
<tr>
<td>9*</td>
<td>M</td>
<td>+</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>Acne</td>
<td>+</td>
<td>0</td>
<td>0</td>
</tr>
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<tr>
<td>32†</td>
<td>M</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Nil else</td>
<td>+</td>
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<td></td>
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*Inferior vena caval thrombosis also.
†Resected aneurysm of popliteal artery.
MU and GU = mouth and genital ulceration.
of the deep veins of the legs extending in one man to the inferior vena cava. The eighth patient, also a young man, had undergone a successful resection of a popliteal artery aneurysm. It was not possible to relate this complaint to the other features of the disease or to the severity of the disorder (Table 8).

GASTROINTESTINAL INVOLVEMENT
This occurred in 3 cases. (1) One young woman had Crohn's disease proven on resection (Case 22). History. A small Pakistani woman of approximately 33 years of age, resident in England for 5 years. She had had mouth and genital ulcers since the age of 30, and 3 years before these symptoms (i.e. in 1966) had complained of erythema nodosum and a polyarthritis of the elbows, left wrist, both shoulders, and knees. X-rays had shown no bony lesions. Chest x-ray was clear. In 1968 she was reported to have had an eosinophilia of 13%; in 1969 she underwent resection of the small intestine for diarrhoea, the barium meal and follow-through having been reported as normal. Pathology report stated, 'A loop of small intestine 25 cm long with a thickened mucosa and wall about the middle. On either side the bowel looks normal. The thickened lesion shows shallow ulcers. Mesentery at this part is hard. Histology shows that the mesentery contains bowel contents and contains a dense, acute, neutrophil infiltrate. Perforation. The bowel shows epitheloid granulomatous lesions with giant cells and transmural infiltrate in places. No tubercle bacilli seen by special stain. The findings are consistent with Crohn's disease.'

After resection joint symptoms worsened. She did not complain of back pain at any time; there was a negative history of sepsis on skin puncture and no other relevant history. Examination showed a single deep slit-like mouth ulcer, absent right labium majorum without scarring or ulceration, reduced forward flexion of the lumbar spine, minimal limitation of elbow extension, tenderness of both sternoclavicular joints, and a flexor tenosynovitis of the left middle finger. Latex and sheep cell agglutination tests for rheumatoid factor were negative, immunoglobulins were high: IgG 17·0 g/l, IgA 4·1 g/l, IgM 2·1 g/l. No x-rays of the sacroiliac joints were taken because of the patient's age. HLA antigens were not determined.

(2) A second patient (Case 17) had a resection of the caecum for a nonspecific caecal ulcer. History. A 33-year-old married woman had had frequent mouth ulcers since the birth of her first child when she was 21. Genital ulceration began at the age of 26 years and there had been much marital disturbance with a suicide attempt. The Blobner phenomenon was positive. The patient had never suffered arthritic, neurological, or vascular involvement and was managing well. She was fairly fit at the time of this survey. When seen again 2 years later she had undergone a right hemicolectomy for abdominal pain thought probably due to Crohn's disease. The caecal ulcer found in the specimen was 4 cm in diameter. The base was 'composed of polymorphic oedematous debris, with a deeper fibroelastic reaction infiltration by chronic inflammatory cells of the plasma cell type with a number of eosinophils. The appearance did not suggest Crohn's disease. Two adjacent lymph glands examined showed reactive hyperplasia.'

(3) A third patient (Case 2), a 19-year-old married woman, had had recurrent ulceration either just anterior to the anus or just inside it. Ulcers were associated with pain and responded to steroids as did her severe mouth ulcers, present since the age of 7. One attack of vulval ulceration had left scarring. The left shoulder was clinically normal at interview but had been the site of previous arthritis.

CLINICAL MEASUREMENTS
Reduced chest expansion, forward flexion, and lateral flexion of the lumbar spine were found in very few patients (Table 7). No correlations could be established with other features of Behcet's disorder.

RADIOLOGY
Radiographs of the sacroiliac joints of 17 patients (6 male [2 with uveitis], 11 female) were available. No sacroilitis was found. Abnormal findings indicative of osteoarthrosis were found in the hands of 2 patients and the feet of 2 others.

BLOOD STUDIES
Estimations of the ESR taken at the time of interview (when the disease was not necessarily active) showed that the ESR was raised (>20 mm in the first hour) in 3 patients but the mean level was only 11 mm (range 2–32 mm). No consistent abnormalities of haemoglobin and white cell count were observed.

HLA ANTIGENS (TABLES 9, 10)
Twenty-eight patients were HLA typed using a microlymphocytotoxicity technique for 21 antigens (Terasaki et al., 1964). The group consisted of 12 males and 16 females, the additional members being accounted for by patients seen by the author in the year 1974–75. HLA B5 and HLA B27 were found in increased frequencies in males and in patients of both sexes respectively.

The overall frequency of B5 is not increased in this series but the increased frequency of the antigen in males is comparable with the findings in Japanese
Table 9  HLA frequencies in Behcet's syndrome

<table>
<thead>
<tr>
<th></th>
<th>Male (n = 12)</th>
<th>Female (n = 16)</th>
<th>Controls (n = 613)</th>
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<th>Male (n = 78)</th>
<th>Female (n = 21)</th>
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<tr>
<td>HLA B5</td>
<td>(17·85%)</td>
<td>(0%)</td>
<td>(9·39%)</td>
<td>15</td>
<td>(71·4%)</td>
<td>NT</td>
</tr>
<tr>
<td>HLA B27</td>
<td>(24·99%)</td>
<td>(21%)</td>
<td>(7·02%)</td>
<td>24</td>
<td>(30·8%)</td>
<td>NT</td>
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<td>= not tested</td>
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Table 10  HLA B27 in Behcet's syndrome

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<th>Case no.</th>
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<th>Phlebitis</th>
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</tr>
<tr>
<td>Total</td>
<td>2/6</td>
<td>1/6</td>
<td>0/6</td>
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</tr>
</tbody>
</table>

Behcet's syndrome in 32 patients in Yorkshire  497

and closely, if infrequently, associated with orogenital lesions. Psoriasis and psoriatic arthropathy are found to be associated with backache, sacroiliitis and raised IgA but dissociated from those features seen in severe Behcet's syndrome.

Discussion

A population of 5 million resident in the northern part of England (the former County of Yorkshire) comprising both highly industrial towns and sparsely populated rural areas yielded 32 subjects found to have recurrent mouth ulceration and other features.

Genital ulceration was found in 91%, nondestructive arthritis in 47%, skin lesions in 44%, vasculitis in 25%, and iritis in 12%. These figures are similar to those for Behcet's disease in Japan given by Oshima et al. (1963). Strachan and Wigzell (1963) working from a different viewpoint found arthritis in all cases but their frequencies of oral and genital ulceration were somewhat lower and almost identical. Cooper and Penny (1974) and Mason and Barnes (1969) found mouth ulceration in all their patients, with genital ulceration in three-quarters. Even those patients of our series who did not have true genital ulceration reported septic lesions in the area so that there is evidence to support the claim that the diagnosis of Behcet's syndrome should not be considered unless bipolar disease (mouth and genital ulceration) is present. One might keep an open mind about prepubertal children: some of these present with gross mouth ulceration and only later develop genital and other features.

While the series as a whole shows the features enumerated by Behcet and features such as erythema nodosum, vasculitis, and eye lesions are shown by Q-analysis to be closely connected with mouth ulceration, only 3 of our patients would satisfy Behcet's criteria for the triad (mouth ulceration, genital ulceration, and lesions of the anterior chamber of the eye). 21 patients satisfy Mason and Barnes' (1969) definite criteria (without using the family history and cardiovascular lesions). All

patients with Behcet's syndrome (Ohno et al., 1973) (for males $\chi^2 = 10·14$; $P = 0·0015$). There is no significant sex difference in the frequency of B27 in Leeds patients with Behcet's (no figures are available for their Japanese counterparts), the overall frequency being significantly increased compared with controls ($\chi^2 = 9·68$; $P = 0·0019$). This remains significant ($P = 0·42$) even when the original P value is multiplied by 21 to take account of the number of antigens tested (Svejgaard et al., 1974).

IMMUNOGLOBULINS
IgG was normal in all subjects, but IgA was raised in 8 and IgM in 18. Rheumatoid factor was positive to a titre of 1 in 256 in one subject and the latex fixation was positive in a total of 6. While indirect immunofluorescence was positive in 11 cases no one system was predominant.

Q-ANALYSIS OF DATA
Data on the patients were compared with those of relatives and spouses. 32 probands and 21 relatives and no spouses of a total of 22 persons surveyed gave a history of mouth ulceration, but in only 32 probands and 10 relatives was the ulceration evaluated as significant. Genital ulceration was closely linked to mouth ulceration in a way which contrasted with latex fixation testing. This was positive in 31 of the 226 persons in the survey but was only weakly connected with orogenital ulceration both in the total analysis and in the analysis of the patients' file. Erythema nodosum, a positive skin test, acne, and vasculitis emerge from analysis as genuinely
patients except one had two of their four major
criteria; this patient had a doubtful history of genital
ulceration (recorded as negative) and thrombo-
phlebitis.
Using this last level of certainty the prevalence of
known Behcet's syndrome in Yorkshire is 0·064 in
10 000. This is probably representative of the
frequency of the condition throughout the United
Kingdom and differs markedly from the figures of
Aoki et al. (1971). In the remote Hokkaido district
off the mainland of Japan they found a prevalence of
1 in 10 000. Of a total of 353 patients, stomatitis was
found in 179, genital ulceration in 107, and serious
ocular lesions in 284. The incidence of the disease
there has apparently risen progressively since 1962,
paralleling a global increase in reported cases:
Strachan and Wigzell (1963) listed a total of 45
patients in the world literature from 1932-61
whereas Chajek and Fainaru (1975) gave a total of
724 cases.
Lumbar pain is not a prominent feature of this
series and is insignificant beside the other manifesta-
tions. Sacroiliitis was not seen in the 17 patients
x-rayed. Hill (1969) questioned whether sacro-
ilitis was part of Behcet's syndrome, and Cooper
and Penny (1974) mentioned a case of 'clinical
sacroiliitis'; the matter is of importance in relating
the disease to sacroiliitis in the known seronegative
polyarthritides and in turn to HLA B27 or HLA
antigens.
While there is a modestly raised incidence of B27
in this series, the level being similar to that found in
psoriasis, the frequency of the antigen is much below
that found in ankylosing spondylitis and even in
Reiter's disease. The British figures confirm the only
reported Japanese series in showing an increased
incidence of A5 although this rise has been found in
males only in the Leeds survey, and its significance
awaits further investigation.

The characteristics of the arthritis and arthralgia
in this series are consistent with those quoted by
Mason and Barnes. Asymmetrical large joint
involvement is relatively common; arthralgia may be
episodic but rarely migratory although several joints
may be involved from time to time. During exacerba-
tions synovial fluid complement is raised and a
polymorphonuclear leucocytosis is present (Zizic
and Stevens, 1975).

Immunoglobulins M and A were raised in this
series. Immunoglobulin A is also raised in psoriasis
(and was in our 1 patient who also had psoriasis).
Tests for antinuclear factor were consistently
negative and although indirect immunofluorescence
tests were sometimes positive no definite pattern
emerged. Changes documented would suggest that
autoimmunity involving the tissues studied is not of
great importance in the pathogenesis of Behcet's
disease. Immunofluorescence against fetal oral
mucosa may be more relevant (Lechner, 1967), or
these mechanisms may perhaps be brought into
action by other as yet unknown factors converging
on the vasculitis which is commonly observed
(France et al., 1951; Enoch et al., 1968).

Behcet himself drew attention to the frequency of
sepsis and Blobner (1937) reported on the diagnostic
value of sepsis occurring at the site of skin puncture.
No false positives were found in the relatives and
spouses of our patients, but a positive skin test had
no prognostic value. While Sezer (1953) reported a
transmissible virus his work has not so far been
repeatable by others and the aetiological importance
of infection (viral or bacterial) remains debatable.
The studies of Wright (1965) and Wright and
Watkinson, (1965) have shown linkages between
bowel disorder such as ulcerative colitis and Crohn's
disease and ankylosing spondylitis, and between
ankylosing spondylitis and psoriasis and Reiter's
disease. There have been several reports of ulcerative
colitis occurring in the course of Behcet's disease
(Böe et al., 1958; Empey and Hale, 1972) and it is
important to differentiate these cases from those in
which Behcet's disease itself has resulted in gastro-
testinal ulcers or other abnormalities. 3 patients
had gastrointestinal involvement in the present
series but one patient had proven Crohn's disease.
The expected coincidence of Crohn's disease and
Behcet's would be 0·00576 per million (based on
Evans and Acheson (1965) which gives the Oxford
Cumulative Prevalence as 9±1·4 per 100 000).
Although the likelihood of the two occurring
together is remote and the possibility exists that they
are associated, the question whether certain diseases
are linked to Behcet's disease can only be resolved
by further studies and perhaps by putting all UK
patients on the ARC register. This should allow
something more meaningful to be said about the
aetiology of Behcet's hinted at by the changes in
immunoglobulins, raised incidence of HLA B27
and B5, the vasculitis, and other clinical features.
It is hoped that the family study of which this report
is part will also be of value.

It is important to realize these patients are at risk
from marital breakdown and self-destruction.
Epstein et al. (1970) document 10 cases with severe
character disorders and believe that these were of
both aetiological and secondary types. None of our
10 patients with neurotic problems had severe
neurological involvement and were in greater
danger from their psychiatric problems than from
the physical effects of Behcet's syndrome. Sircus
et al. (1957) found a significant number of patients
with depression and anxiety among subjects with
focal oral ulceration. This is particularly interesting as this common diagnosis may be the fertile ground on which Behcet’s syndrome grows.

I am greatly indebted to Professor V. Wright, Mrs. J. Moll, Guthrie Scott, S. M. Rajah, G. Bevans, D. J. Lintott, I. Haslock, C. Eastmorn, Mrs. B. Gordon, Mrs. J. Packter, Mr. Brian Bentley, and Sister M. Binden of the Leeds Teaching Hospitals and University; and to Mr. R. Atkin and Mr. J. Johnson of the Department of Mathematics, University of Essex. This work would not have been possible without their help and that of many others, including the families interviewed.

References


Behcet's syndrome in 32 patients in Yorkshire.

M A Chamberlain

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