Spondylodiscitis and ankylosing spondylitis: HLA typing and nosological implications

V. MODENA*, N. MIGONE†, V. DANEO*, A. O. CARBONARA†, S. DI VITTORIO†, AND M. VIARA**

SUMMARY Nine cases from among 64 patients with ankylosing spondylitis (AS) are described. In addition to bilateral sacroiliitis these cases had a peculiar type of spondylodiscitis characterised by quite diffuse and marked sclerosis of multiple vertebral bodies, with only minimal erosions of the adjacent vertebral plates while classical syndesmophytosis was absent. The antigen HLA-B27 was found only in 1 of these 9 cases. This type of spondylodiscitis could discriminate among all the patients with AS a subgroup with a peculiar clinical pattern and a probably distinctive pathological mechanism.

Erosive sclerotic lesions of the vertebral bodies surrounding a disc—spondylodiscitis (SD)—have been reported in between 1 and 28% of patients suffering from ankylosing spondylitis (AS) (Forestier et al., 1956; Jaqueline, 1956; Schulitz, 1969; Daneo and Di Vittorio, 1970; Cawley et al., 1972; Doury et al., 1975). Different methods in selecting the patients, different extent of spinal survey, and lack of agreed diagnostic criteria for spondylodiscitis may account for this wide variation in prevalence.

There is no agreement about the origin of these lesions, some authors favouring traumatic (Kanefield et al., 1969; Rivels and Freiberger, 1969; Cawley et al., 1972; Gougeon et al., 1977), others inflammatory mechanisms (Romanus and Ydén 1952; Coste et al., 1963; Little et al., 1974). In the most destructive lesions, often occurring in the late stages of AS at only one vertebral metamer, namely at the hinge-point between two ankylosed spinal segments, the traumatic mechanism is likely.

A peculiar feature of a different SD in subjects diagnosed as having AS was described by Daneo and Di Vittorio (1970). This type is characterised by quite diffuse and marked sclerosis of multiple vertebral bodies with only minimal erosions of the adjacent vertebral plates; it has been observed in patients with bilateral definite sacroiliitis and symptoms or signs referred to the spine, in absence of clear syndesmophytosis.

The HLA typing of patients affected by AS complicated by this latter feature of spondylodiscitis is reported in the present study. The low frequency of the B27 antigen together with peculiar clinical and radiological findings are discussed.

Patients and methods

From a series of 64 patients fulfilling the diagnostic criteria for AS (Gofton, 1968) 9 subjects (14%) had radiological signs of SD. These 9 patients, together with their available first-degree relatives (>16 years of age)—2 parents, 3 sibs, and 3 children—underwent clinical and radiological investigation. Information on the health status of other 38 first-degree relatives was also collected by means of a direct interview with the proband. A total of 17 subjects were typed for HLA-A, B, C specificities as described previously (Daneo et al., 1977). A panel of 1428 healthy individuals constituted the control for HLA antigen frequency.

Results

All the 9 patients were males, with a mean age of 39.2±11.3 years (Table). The first clinical symptom appeared at a mean age of 21.7±5.5 years and was represented by persistent lumbar pain, followed later by gluteal complaints. The pelvic x-rays showed severe bilateral sacroiliitis (grade 3-4).
Spondylodiscitis and ankylosing spondylitis: HLA typing and nosological implications

Table Main clinical, radiological, and HLA data of 64 AS patients, according to the presence or absence of spondylodiscitis (SD)

<table>
<thead>
<tr>
<th></th>
<th>With SD (n=9)</th>
<th>Without SD (n=55)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age</td>
<td>39.2±11.3</td>
<td>40.2±12</td>
</tr>
<tr>
<td>Sex (males/females)</td>
<td>9/0</td>
<td>52/3</td>
</tr>
<tr>
<td>Initial complaint</td>
<td>Lumbar</td>
<td>Gluteal</td>
</tr>
<tr>
<td>Mean age at onset</td>
<td>21.7±5.5</td>
<td>26.9±8.5*</td>
</tr>
<tr>
<td>Bilateral sacroiliitis</td>
<td>9 (100%)</td>
<td>55 (100%)</td>
</tr>
<tr>
<td>Syndesmophytosis</td>
<td>1 (11.1%)</td>
<td>47 (85.4%)†</td>
</tr>
<tr>
<td>B27 antigen positivity</td>
<td>1 (11.1%)</td>
<td>48 (87.3%)‡</td>
</tr>
</tbody>
</table>

*P=0.08 (Student’s t test). †P=5.5 (Fisher’s exact test). ‡P=2.5 (Fisher’s exact test).

According to Goffin (1968) in all 9 cases. The spondylodiscitic lesions were characterised by diffuse and marked sclerosis of vertebral bodies, with minimal erosions of the adjacent vertebral plates (Fig.). The findings were at 3 or more levels in 5 cases, at 2 levels in 3 cases and at 1 level in 1. However, the absence of classical features of syndesmophytosis, except 1 case (B27-negative) is worth noting. In this patient the process was confined to one vertebral body and to an initial stage. Moreover voluminous intervertebral bony bridges recalling the osteophytic and not the syndesmophytic-like pattern were documented in 2 subjects aged 33 and 41, respectively.

The HLA typing showed that 8 out of 9 were B27-negative and only 1 B27-positive. No other HLA specificities were significantly different when compared with the antigen frequencies of the control population. Nevertheless, out of 9 cases 6 carried the HLA-A2 and 4 the HLA-B5 antigen.

Preliminary family analysis showed only 2 affected first-degree relatives; 2 brothers with sacroiliitis in the family of the B27-positive proband.

Discussion

Spondylodiscitic lesions have been documented in patients affected by AS. Some authors suggest that these vertebral involvements could be a possible late complication of classical AS, following a known or suspected trauma upon an ankylosed spine. Others think that these unusual signs might be related to the same inflammatory mechanism of the primary disease.

Recently, Danco and Di Vittorio (1970) and Little et al. (1974) have stressed the precocious occurrence in some AS patients of primary spondylodiscitic lesions at multiple levels, in the absence of classical spinal syndesmophytosis. These latter findings, together with the tendency for voluminous intervertebral bony bridges to form, have suggested a different pathogenetic mechanism for this particular type of spondylodiscitis. The results obtained in this study seem to confirm these previous observations. In our 9 patients some clinical and radiological findings display peculiar features which distinguish them from classical AS (Table).

The main differences lie in the radiological aspect of the spine, which is characterised apart from SD, by the absence of typical syndesmophytosis and, eventually, by a precocious tendency to show voluminous osteophytic-like bony bridges. These bony bridges recall those occurring in infectious spondylodiscitis.

A more striking difference is seen in the HLA typing which shows the absence in our 9 spondylodiscitic cases of the ‘expected’ increase of the B27
antigen frequency; in fact only 1 out of 9 (11.1\%) was B27-positive. Furthermore, the preliminary findings of no secondary AS or sacroilits among the relatives of 8 B27-negative AS probands seem also to indicate a different genetic susceptibility from classical AS. On the other hand, the observation of 2 brothers affected by sacroilits in the family of the B27-positive SD patient is in a good agreement with the high familial occurrence in families of B27-positive AS patients (Contu et al., 1977; Daneo et al., 1977; Kidd et al., 1977).

The absence of B27 in our patients formerly included in the AS category is difficult to explain because we have no definitive data on the pathogenesis of the disease. One hypothesis could be that the same environmental stimulus which triggers the classic and complete form of AS in a B27-positive case, might develop in some B27-negative individuals, as well as the sacroilits, some peculiar spinal lesions with a spondylodiscitic-like pattern. This view is supported by the finding in 3 out of 7 B27-negative AS patients (aged 34, 38, and 47) of voluminous intervertebral bony bridges, possibly suggesting a previous SD involvement. Whether or not this peculiar SD feature represent a major distinctive marker in B27-negative AS, further studies on larger series of patients lacking B27 antigen are clearly needed.

In conclusion, these data show that a type of spondylodiscitis characterised by early onset, involvement of multiple vertebral bodies, predominance of sclerosis upon erosion, tendency to show osteophytic-like bony bridges at the spine rather than typical syndesmophytosis is almost solely restricted to B27-negative AS patients.

It seems possible that a different prevalence of this type of SD accounts partly for the reported deviation of the B27 positivity in some AS studies.

References


