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HLA B27 and the genetics of ankylosing spondylitis

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SUMMARY One hundred and twenty-eight of 145 patients with ankylosing spondylitis (AS) were found to be HLA B27 positive. Five patients had evidence of a sero-negative peripheral arthritis resembling peripheral psoriatic arthritis and 3 of these were B27 negative. One further B27 negative patient had a sister with ankylosing spondylitis and ulcerative colitis and a mother with ulcerative colitis. There was evidence of a somewhat later age of onset of symptoms in B27 negative patients. These findings are interpreted as suggesting some degree of clinical and genetic heterogeneity in ankylosing spondylitis with genes for psoriasis and inflammatory bowel disease being important in some individuals, particularly those who are B27 negative.

Twenty-five first-degree relatives with ankylosing spondylitis were all B27 positive. The only instance of disassociation of B27 and spondylitis in a family was where the proband had ulcerative colitis as well as spondylitis. Of 13 B27 positive fathers 3 could be diagnosed as having definite ankylosing spondylitis (23%). These findings are thought to provide evidence against the concept that the gene for ankylosing spondylitis is not B27 but a closely linked gene and favour the occurrence of an environmental event affecting approximately one-fifth of B27 positive males to result in disease.

The finding by Brewerton et al. (1973) and Schlossstein et al. (1973) of a markedly increased frequency of HLA B27 in patients with ankylosing spondylitis raises important questions concerning the genetics and pathogenesis of this disorder. The aims of the present study were to examine the relationship of B27 to ankylosing spondylitis by studying a series of patients to determine (1) if there was evidence of clinical and genetic heterogeneity in ankylosing spondylitis, (2) if dissociation of ankylosing spondylitis from B27 occurs in families, and (3) the prevalence of ankylosing spondylitis in first-degree relatives of B27 positive probands.

Patients and methods

Patients were consecutively included in the series after clinical and radiological assessment resulting in a diagnosis of definite ankylosing spondylitis as judged by the New York criteria (Bennett and Wood, 1968). Patients who were found to have psoriasis or inflammatory bowel disease or patients with spondylitis who had been ascertained through a study of patients with uveitis were excluded. Histocompatibility antigen typing was performed in these patients and in 451 controls who were blood donors or members of staff, using a standard microlymphocytotoxicity technique (Terasaki and McClelland, 1964).

FAMILY STUDIES

These can be divided into 3 separate studies. The first was selective, in which relatives suspected of having a relevant rheumatic disease from the history given by the proband were examined clinically and radiologically and HLA typed.

The second was a clinical, radiological, and HLA typing study of parents of 20 HLA B27 positive probands, where both parents were alive and accessible. In addition, the parents of 2 further HLA B27 positive patients with ankylosing spondylitis ascertained because of uveitis were included. Parents were chosen for this study as all were over the age of 45 years and it is unlikely that evidence of ankylosing spondylitis will appear for the first time after this age.

Thirdly, 3 identical twins of patients with ankylosing spondylitis were examined, and the results have been previously reported (Eastmond and Woodrow, 1977).

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Results

HLA TYPING

One hundred and twenty-eight of the 145 probands (88.3%) were HLA B27 positive, the control frequency being 8.97%. Twenty-two of the patients were female of whom 1 (4.5%) was B27 negative compared with 16 of the 122 males (13%). A history of anterior uveitis was obtained in 22 patients (15.9%) of whom 16 were male and all were B27 positive with the exception of one male patient.

PATIENTS WITH PERIPHERAL ARTHROPATHY

One B27 positive male patient had, in addition to ankylosing spondylitis, sero-positive erosive peripheral rheumatoid arthritis with a histologically confirmed rheumatoid nodule over the left olecranon process.

Two male B27 positive patients had a history of previous Reiter's disease occurring several years before the onset of the symptoms of spondylitis.

Five patients had a sero-negative peripheral small joint arthropathy with features similar to those seen in patients with peripheral psoriatic arthropathy, 2 being B27 positive and 3 negative. The 2 B27 positive patients both gave a history of 'sausage toes' and 1 of these, who previously had onycholysis of some of the finger nails but no skin lesions, has more recently developed a small skin lesion on the right knee suggestive of psoriasis.

The 3 B27 negative patients in this category are of considerable interest. One had had episodes of 'sausage fingers' and some swelling of proximal interphalangeal joints and later developed swan-neck deformity of several fingers with minimal erosive changes radiologically. The second patient has had swelling and tenderness of interphalangeal joints of the toes and his father has psoriasis with similar changes in the toes and a proximal interphalangeal joint arthropathy in the hands but no radiological sacroiliitis. The third patient has had an asymmetrical distal and proximal interphalangeal joint arthropathy of the hand with erosive changes in the clinically affected joints.

Two B27 positive female patients have had a mild peripheral small joint oligoarthritis with no distinctive features in respect of joints affected or clinical features of the joint involvement.

HLA B27 NEGATIVE PATIENTS

Three of these 17 patients are documented above. Two other patients in this group are of interest. One male developed symptoms of ankylosing spondylitis at the age of 32 years and at the age of 62 years developed an acute synovitis of wrists, metacarpophalangeal joints, and ankles with associated circinate balanitis, keratoderma blennorrhagicum of the palms and soles, and conjunctivitis but no urethritis or diarrhoea. This episode resolved leaving no residua. The second patient, also male, and whose parents were first cousins had a sister with ankylosing spondylitis and ulcerative colitis and a mother with ulcerative colitis and probable ankylosing spondylitis. (Fig. 1).

The 12 remaining B27 negative patients differed in no way clinically or radiologically from the B27 positive patients. Six of these were of considerable clinical severity with radiological changes in the spine as well as in the sacroiliac joints.

AGE OF ONSET (Fig. 2)

It was possible to determine the approximate age of onset with some reasonable accuracy in 137 patients. The distribution of the age of onset in B27 negative patients was compared with that in the B27 positive patients by the Wilcoxon rank sum test. It was found that the age of clinical onset in B27 negative patients was higher overall than that in the B27 positive patients (P=0.012).

FAMILY STUDIES

Selected families

In the families of 22 probands all of whom were B27 positive there were 2 or more cases of ankylosing spondylitis. Four of these probands were female. All 25 clinically affected relatives of these probands were B27 positive. The only instance where an affected relative of a B27 negative proband was found is shown in Figure 1.

Figure 3 shows a family not included within the main series because the B27 negative proband had ulcerative colitis in addition to ankylosing spondylitis.
Study of parents

Thirteen of the 22 fathers and 12 of the 22 mothers of 22 HLA B27 positive probands were found to be HLA B27 positive. Of these parents, 2 fathers were known to have clinical and radiological ankylosing spondylitis and 1 father had asymptomatic grade 2 radiological sacroiliitis and spondylitis with limited spinal movements and diminished chest expansion. Two fathers had asymptomatic grade 3 radiological sacroiliitis. Thus 3 out of 13 (23.1%) B27 positive fathers could be diagnosed as having ankylosing spondylitis. No clinical or radiological evidence of disease was found in the HLA B27 positive mothers or in the HLA B27 negative parents.

Identical twins

A study of 2 identical twins of probands with ankylosing spondylitis revealed no clinical or radiological evidence of ankylosing spondylitis or related disease. Their ages were 45 and 52 years, respectively. The identical twin aged 59 of a further patient with severe ankylosing spondylitis had a history of 1 episode of acute anterior uveitis and radiologically had a unilateral grade 3 sacroiliitis but no rheumatic symptoms (Eastmond and Woodrow, 1977).

Discussion

The association between ankylosing spondylitis and HLA B27 raises two main and related questions. The first is whether the major gene underlying susceptibility to the disease (the AS gene) is B27 itself or whether it is not B27 but a gene which is at a locus very close to the HLA B locus and which is in very strong linkage disequilibrium with B27. The second and related question is whether the spondylitis in B27 negative individuals is genetically the same as in B27 positive subjects, i.e. that the AS gene is the same for both groups and is therefore not B27.

B27 ITSELF VERSUS LINKED AS GENE

A strong possibility in regard to most HLA and disease associations is that the disease susceptibility gene may not be the particular HLA A or B locus genes found to be associated with the disease but another genes at closely linked loci in linkage disequilibrium with the A and B locus genes. However, the relative risk for ankylosing spondylitis in B27 positive individuals when estimated from combined data 28.8 (Woodrow, 1977), this being considerably higher than for any other HLA and disease association.

If the HLA linked AS gene were not B27 itself two findings might be expected. The first is that aggregation of B27 negative individuals with spondylitis should occur in families to the same degree as observed in the case of B27 positive patients. So far...
no pair of B27 negative first-degree relatives with spondylitis appears to have been documented. The only instance we have observed is the family shown in Fig. 1 in which ulcerative colitis was present in 1 of the 2 affected B27 negative sibs and in their mother. It is of interest that the parents in this family were related, suggesting the possibility of a recessive trait predisposing to spondylitis. Another interesting family in which a B27 negative spondylitic was the offspring of a consanguinous marriage was reported by Van der Linden et al., (1975).

Secondly, one might expect to see examples of disassociation of B27 from ankylosing spondylitis within a family, ie, evidence of recombination between the B locus and the supposed locus for the AS gene. Two families have been reported (Dick et al., 1975; Strosberg et al., 1975) in which apparent disassociation of spondylitis from B27 has occurred. In these instances genetic recombination would be the explanation if it were certain that other, non-HLA linked genes predisposing to spondylitis were not segregating in the families.

The only occasion on which this phenomenon was observed in the present study was in a family shown in Fig. 3 where the B27 negative patient with spondylitis also had ulcerative colitis.

The demonstration of genetic recombination classically depends on the certain identification of the presence or absence of relevant segregating genes and thus it is relatively easy for traits showing simple Mendelian inheritance, ie, the presence of a particular phenotype must imply the presence of one particular gene. This does not necessarily apply to conditions with a possible heterogeneous genetic basis and where the trait does not invariably appear in the genetically predisposed.

Further information on this point may be obtained by studying the prevalence of ankylosing spondylitis in B27 positive persons in the general population compared with B27 positive relatives of ankylosing spondylitis probands. Evidence has been produced that something of the order of 20% of B27 positive individuals develop ankylosing spondylitis in some degree (Calin and Fries, 1975; Truong et al., 1975; Cohen et al., 1976). It is of interest to compare this figure with that for the incidence of disease in B27 positive first-degree relatives of B27 positive probands. If indeed the 20% incidence claimed is the true figure for the general population it could arise in three main ways. The first is that the AS gene is B27 and a separate gene segregating independently of the HLA system and present in homozygous or heterozygous state (depending on whether the effect is as a recessive or dominant trait) is also necessary to produce the complete genotype for the disease. If this were true one would expect that, for a dominant interacting gene, 50% of first-degree B27 positive relatives would develop spondylitis and, for a recessive trait, approximately 25% of B27 positive sibs and a considerably lower frequency of parents would develop disease. Secondly, B27 might be the AS gene and an environmental factor randomly affecting 20% of the population is necessary for the disease to develop. This would result in a similar prevalence of disease in B27 positive relatives of probands as in B27 positive persons in the general population, ie 20%, unless these relatives are more liable to be exposed to an important environmental agent. The third possibility is that the AS gene is not B27 but is present on approximately 20% of chromosomes which have the B27 gene. In this case all B27 positive first-degree relatives would be expected to have this gene and to develop disease to some degree.

The finding of ankylosing spondylitis in 23.1% of B27 positive fathers in the present study may be compared with the figure given above for the general population. The fact that these are of a similar order is strongly against a linked AS gene on 20% of B27 positive chromosomes, and is most in keeping with the result expected with the second proposal of a random environmental factor affecting 20% of the population. However, this approach requires further investigation using identical criteria for the study of the general population and the families. Discordance in identical twins is also supportive evidence for environmental factors affecting the prevalence of disease in genetically predisposed individuals.

The absence of clinical and radiological abnormalities in any of the B27 positive mothers studied so far supports previous general experience of the sex distribution of the disease and is difficult to reconcile with the findings of Calin and Fries (1975) who in their study of a population of B27 positive individuals found approximately equal numbers of males and females with evidence of spondylitis.

**B27 POSITIVE SPONDYLITIS versus B27 NEGATIVE SPONDYLITIS**

In regard to the previous discussion it is of some interest to know if there is evidence for heterogeneity within the disease ankylosing spondylitis, as this may have a bearing on the way we may attempt to determine the nature of the AS gene.

**CLINICAL EVIDENCE**

It has been suggested that the disease takes a milder and more localised course in B27 negative patients. Thus Möller and Olhagen (1975) found all 66 patients with radiologically demonstrable syndesmophytes to be B27 positive, in contrast with 70% of 60 patients
with sacroiliitis as the only radiological manifestation. Feldmann et al. (1975) found 6 B27 negative patients amongst 25 with milder disease and only 1 of 25 with moderate or severe disease. However, Van den Berg-Loonen et al. (1977) found 4 of 20 patients with spinal involvement to be B27 negative and Jeannet et al. (1975) found 3 B27 negative patients among 14 females with radiological involvement of the spine. In the present series the 17 B27 negative patients had clinical and radiological involvement as severe as was found in the B27 positive patients. One possibility is that patients with sacroiliitis only are aetiological more heterogeneous than those with spinal changes and it is of interest that Dekker-Saeyes et al. (1978) described 11 patients with ulcerative colitis who had asymptomatic radiological sacroiliitis and of these only 1 was B27 positive.

With regard to peripheral arthritis, 3 of the B27 negative patients in the present study had an arthropathy with features strongly suggestive of psoriatic arthropathy but in the absence of psoriasis. It has been shown that patients with psoriatic peripheral arthritis and ankylosing spondylitis are less frequently B27 positive than patients with ankylosing spondylitis alone (reviewed in Woodrow, 1977). The implication is that genes for psoriasis may be playing a role in the pathogenesis of spondylitis in these cases, a conclusion also drawn from a study of the arthropathies occurring in patients with psoriasis (Eastmond and Woodrow, 1977).

The fact that B27 negative patients tend to have a somewhat higher age of onset of symptoms supports the thesis of aetiological heterogeneity between the two groups.

As mentioned above, no examples of ankylosing spondylitis occurring in 2 B27 negative relatives has been reported previously. The only example seen by us was a family in which there could be genes for ulcerative colitis predisposing to the development of ankylosing spondylitis. This explanation is supported by the fact that patients with ulcerative colitis and ankylosing spondylitis are less frequently B27 positive than those with ankylosing spondylitis alone (Brewerton et al., 1974). The family shown in Fig. 3 also gives credence to this argument. The brother without ulcerative colitis could be considered as dependent upon being B27 positive in order to develop ankylosing spondylitis whereas the proband with ulcerative colitis was able to develop equally severe spinal disease in the absence of B27.

**Conclusions**

(1) Present evidence favours the gene for ankylosing spondylitis being the HLA B27 gene itself and the results of present study would be against the linked-gene hypothesis.

(2) There is some evidence that ankylosing spondylitis is genetically heterogeneous and that in HLA B27 negative persons it differs from the disease in HLA B27 positive persons by (a) having a later mean age of onset, (b) having, in some cases, features of psoriatic peripheral arthritis, and (c) having, in some cases, evidence for genes for ulcerative colitis predisposing to spinal disease.

(3) The present study does not support the suggestion that ankylosing spondylitis is less severe in HLA B27 negative patients than HLA B27 positive ones, when no associated disease is present.

**Association with other arthropathies**

The single case of sero-positive rheumatoid arthritis is considered to represent the coincident occurrence of the separate conditions in the same individual.

The two instances where Reiter's disease had occurred before the onset of symptoms of spondylitis probably represent the occurrence of two diseases provoked in the same genetically predisposed individuals by different environmental agents. Instances where Reiter's disease occurred in patients who already had ankylosing spondylitis have been previously reported (Woodrow et al., 1974).

**References**


