Experimental autoimmune arthritis in mice. I. Homologous type II collagen is responsible for self-perpetuating chronic polyarthritis

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SUMMARY Immunisation with heterologous type II collagen (CII) induces arthritis in mice of the DBA/1 strain, which is genetically susceptible to this disease. To develop an experimental model of autoimmunity more adequate for the study of human rheumatoid arthritis (RA), DBA/1 mice were injected with 100 μg of native CII that had been purified from mouse xiphoid cartilage. About six weeks later the animals developed a chronic progressive polyarthritis involving the four paws but mainly confined to interphalangeal and metatarsophalangeal joints. The evolution of the disease fluctuated between remissions and exacerbations. The initial lesions assessed by clinical observations were more severe when the disease occurred early than in the case of late onset. Interestingly, the incidence of arthritis was clearly preponderant in males, and, moreover, the few female mice which developed arthritis had mild disease states with lower arthritic scores than the males. Varying levels of autoantibodies against mouse CII were found in the sera of immunised animals, regardless of the development of arthritis. These data indicate that the injection of homologous CII into mice caused a polyarthritis that is clinically closer to the human RA than the disease induced with heterologous CII and therefore will represent a useful tool for the study of the self-perpetuating mechanisms that characterise RA.

Key words: murine type II collagen, rheumatoid arthritis, sex linkage, anticollagen autoantibodies.

Rheumatoid arthritis (RA) is characterised by a chronic proliferative synovitis of diarthrodial joints that eventually leads to articular cartilage and bone erosion. The precise nature of the immune reactions that take place in the synovial tissue in RA is poorly understood. It has been postulated that the massive infiltrates of helper T cells in close contact with HLR-DR+ synovial stellate cells are mutually stimulatory, supporting the existence of a local self-perpetuating activation.1,2 Moreover, the susceptibility to this autoimmune disease is regulated by major histocompatibility complex genes. Thus the risk of developing RA is associated with DR4 antigens.3,4

In the past, numerous models of experimentally induced arthritis were developed using adjuvants, antigens, or type II collagen (CII) as immunising agents, with a view to analysing the pathogenesis of RA. Since patients suffering from chronic inflammatory arthritis have humoral5-13 and cellular10-12,16 sensitivities to collagen, collagen induced arthritis (CIA) may have relevance to the human disease. First described in rats17,18 and then in mice,19-20 the collagen induced experimental model shares a number of clinical and histological features with RA.21 Likewise, susceptibility to CIA is under genetic control in both rats22,23 and mice.20,24 Moreover, the formation of a synovial pannus and the accompanying articular cartilage erosion are two morphological manifestations of CIA. This experimental disease, however, is provoked by injection of a xenogeneic protein probably carrying species specific determinants that may be immunogenic and contribute to the development of the arthritis.
Therefore, we attempted to induce an experimental autoimmune arthritis (EAA) that would more closely mimic the human RA by injecting native CII of murine origin into the susceptible DBA/1 mice.

**Materials and methods**

**MICE**

DBA/1 mice aged 6 to 17 weeks were obtained from the Institut Pasteur breeding centre (Paris, France). Animals matched for age and sex were used in all experiments.

**Cartilage Collagens**

Mouse CII was prepared from the xiphoid appendix. After careful dissection to eliminate fatty tissue and avoid bone contamination the xiphoid cartilages were pulsed in liquid nitrogen and the powder obtained was then washed at 4°C with 0·05 M TRIS/2 M MgCl₂ buffer, pH 7·4, and further rinsed twice with distilled water. The residue was solubilised in 0·5 M acetic acid (adjusted to pH 2·5 by the addition of formic acid), and digestion with pepsin (Sigma, St Louis, Mo, USA) proceeded for 72 h at 4°C. After neutralisation and then centrifugation at 20 000 g for 20 min the undigested residue was treated twice with pepsin under identical conditions. Supernatants from the three extractions were dialysed against 0·05 M TRIS/0·2 M NaCl, pH 7·6, and passed through a column of diethylaminoethyl (DEAE) cellulose. Collagen was then eluted with the same buffer, precipitated by addition of 3 M NaCl, solubilised with 0·5 M acetic acid, and dialysed against 0·01 M Na₂HPO₄. The precipitate was dissolved in acetic acid, and collagen was again precipitated by adding 1 M NaCl, redissolved in acetic acid, dialysed against 0·1 M acetic acid, and finally lyophilised. The purity of murine CII preparations was verified by the presence of a single band after electrophoresis on polyacrylamide gel.

Native bovine type II collagen was kindly provided by D Herbage (CERAD, Lyon, France).

**Immunisation Procedures**

Lyophilised native CII was dissolved in 0·1 M acetic acid at 4°C and this solution was emulsified with an equal volume of complete Freund’s adjuvant (CFA; Gibco Laboratories, New York, USA). Two different immunisation protocols were used: each group was composed of 48 mice including males (M) and females (F). In group I, 14 mice (6M, 8F) were primed intradermally with 100 μg of native mouse CII (in 200 μl) at four to six different sites on the back and were challenged 21 days later with 100 μg of native mouse CII in acetic acid, injected intraperitoneally. As a positive control, 8 M and 6 M received the same doses of native bovine CII using identical immunisation procedures. As negative controls, 10 mice (5M, 5F) were primed with CFA alone and challenged with acetic acid, and 10 other mice (5M, 5F) remained untreated. Group II included 13M and 14F injected intradermally with 100 μg of native mouse CII in CFA (in 100 μl) into the right hind foot pad. Females, but not males, were boosted intradermally 14 days later with an additional 50 μg of mouse CII in incomplete Freund’s adjuvant (IFA; Gibco). As controls for the males, 5M were primed once with bovine CII and 11M received CFA alone. Following the same immunisation protocol as that used for the females injected with mouse CII, 5F were injected first with CFA alone and 14 days later with IFA alone.

**Arthritis Evaluation**

Animals from both groups were observed daily for the onset of arthritis, beginning three weeks after priming until they were killed at nine weeks post-priming for the first group and 24 weeks post-priming for the second group. The date of disease onset was recorded on a daily basis for individual mice. After clinical examination of the non-injected paws (four limbs for group I and three limbs for group II) the severity of the disease was graded by an arthritic index based upon the macroscopic signs observed (erythema, swelling, distortion of the joints) and calculated by adding the scores of each individual joint affected.

**Table 1: Incidence of arthritis induced with native homologous or heterologous type II collagen**

<table>
<thead>
<tr>
<th>Group</th>
<th>Sex</th>
<th>CFA</th>
<th>Mouse CII</th>
<th>Bovine CII</th>
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<td>0</td>
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<td>II*</td>
<td>M</td>
<td>0/11</td>
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* Mice from group I were primed with 100 μg of native CII (or acetic acid alone) emulsified in CFA injected intradermally at four to six sites on the back and were challenged intraperitoneally 21 days later with 100 μg of CII of the same origin as that used for priming (or acetic acid alone).

†Mice from group II were injected with 100 μg of CII (or acetic acid alone) in CFA intradermally into the right hind foot pad. Females, but not males, were boosted intradermally with an additional 50 μg of CII (or acetic acid alone) in IFA 14 days later.

¹ND = not done.
HISTOPATHOLOGY
The animals were killed and the four limbs of all animals (including the non-injected control mice) were dissected out for histological examination. Joints were slowly decalcified for one month and embedded in paraffin blocks. At least five sections (5 μm thick) were cut along a longitudinal axis, mounted, and stained with haematoxylin and eosin before being evaluated blind.

ANTI-CII ANTIBODY MEASUREMENT
Mice were bled by retro-orbital puncture on several occasions during the experiments as reported in the text. The individual serum samples were collected, divided into aliquots, and stored at −70°C until they were tested for antibodies directed against mouse and bovine CII using an enzyme linked immunosorbent assay (ELISA) adapted from Stuart et al. [25] MicroELISA plates (Nunclon, Roskilde, Denmark) were coated overnight at 4°C with 50 μl/well of a 10 μg/ml solution of mouse (or bovine) CII in phosphate buffered saline (PBS). After three washes with 0.1 M TRIS/0.15 M NaCl containing 0.05% Tween 20, pH 7.4 (TRIS-Tween buffer), the remaining protein binding sites on the plastic were blocked by the addition of 0.25% bovine serum albumin (Sigma) in PBS for two hours at 4°C. The plates were then washed three times with TRIS-Tween buffer and 50 μl of the serum to be tested (diluted 1:100) was deposited in duplicate wells. Plates were kept overnight at 4°C, washed three times with TRIS-Tween buffer, then the appropriate dilution of goat antimouse immunoglobulins coupled to alkaline phosphatase (Miles-Yeda Laboratories, Israel) was added to each well. The amount of bound enzyme was measured with p-nitrophenol as the chromatogen substrate in a Hitertek multispec spectrophotometer at 405 nm. The results were expressed as optical density (OD) × 10³ in experimental wells minus the background OD obtained in wells in which the CII coating had been omitted.

Results
INCIDENCE OF ARTHRITIS INDUCED WITH NATIVE HOMOLOGOUS CII
In a first series of experiments mice were primed with 100 μg of native mouse CII (or bovine CII as a positive control) and challenged three weeks later with the same dose of respective antigens, as described in 'Materials and methods'. The administration of homologous CII elicited a high incidence of autoimmune arthritis in DBA/1 mice. Interestingly, the males were more susceptible to the

Fig. 1 (A) Hind paw of a control DBA/1 mouse injected with CFA and (B) that of an arthritic male mouse eight weeks post-immunisation with native mouse CII. There was diffuse soft tissue swelling and intense erythema of the entire paw with prominent synovitis of the knee, the ankle, and all the metatarsophalangeal and interphalangeal joints. Note the markedly increased thickness of the first, fourth, and fifth toes.
induction of EAA regardless of whether they received the homologous or the heterologous protein (Table 1). To confirm the sex linkage of EAA, in the second series of experiments males were immunised only once with native mouse CII, whereas females were injected twice. Despite the booster injection, most of the females remained resistant to the induction of the disease, whereas all the males became arthritic (Table 1). When the results of both groups of mice were combined and analysed by the χ² test the preponderance of EAA was significantly higher for males than for females (p<0.001).

**CLINICAL COURSE OF EAA**

All mice were clinically examined for evaluation of their disease, beginning three weeks post-priming with CII, until nine weeks for group I, and 24 weeks for group II. Control mice injected with CFA alone, or non-injected controls, were simultaneously examined and did not show any macroscopic signs of arthritis. Fig. 1 compares the hind paw of a male mouse that developed a severe autoimmune arthritis with a non-arthritic limb from a control mouse.

As shown in Table 2 the onset of arthritis was significantly delayed when the mice received homologous CII in comparison with immunisation with heterologous CII. Evaluation of the arthritic score showed that the disease severity was not significantly different in the group of male mice injected with mouse CII than in those injected with bovine CII (Table 2). In the latter case the disease developed explosively one month after the administration of the xenogeneic protein and the maximum severity was rapidly attained. After the acute phase, exacerbation of the arthritis did not occur spontaneously. In contrast, the disease provoked by the mouse CII injection was much more progressive and its evolution fluctuated between remissions and exacerbations even after a single injection of CII. Fig. 2 illustrates the development of clinical arthritis in six individual male mice from group II. Typically, interphalangeal and metatarsophalangeal joints were gradually and chronically affected. Acute symptoms generally disappeared rapidly and then recurred several weeks later in the same or other joints. During the quiescent period of the arthritis the affected joints never completely recovered from the inflammation, therefore when the disease relapsed the primarily affected joints showed additional lesions. In comparison with the males, the few females that developed the disease had disparate times of onset but always showed a mild disease form (Table 2).

Interestingly, we found an inverse relationship between the severity of the primary lesions and the
time of disease onset (Fig. 3). Thus the earlier the first macroscopic signs of arthritis appeared, the higher the arthritic scores observed during the initial phase of the disease.

**Histological Studies**

Histological sections were performed at the time of death on the four paws for mice from group I, and on the three non-injected paws for animals from group II. Sections of the limbs that were injected with the antigen were examined separately and were always characterised by a dense inflammatory cell infiltrate beneath the hyperplastic synovial cells, around the capillary blood vessels and often extending more deeply into the adipose tissue. Conversely, in the non-injected paws a continuous spectrum of synovitis was observed, the histological lesions were usually asymmetrical and involved more frequently, and more intensely, the distal rather than the proximal joints. All the affected joints showed synovial proliferation with excessive convolution and increased degrees of multilayering (Fig. 4A). In many cases synovial cells had a greater amount of cytoplasm, became cuboidal, and were dissociated by an intrasynovial oedema (Fig. 4B). Frequently, adjacent to the abnormalities of the synovial lining cells, the sections showed a single layer of flat synovial cells in a different area of the same joint. In some instances the mitosis of synovial cells was accompanied by the deposition of fibrin-like material on the synovium.

The histological lesions observed were consistently milder in mice injected with homologous CII.
than in those given bovine CII. At the time of death the joints from animals immunised with mouse CII showed a sparse focal infiltration of mononuclear cells into the subsynovial area or even deeper into the hypodermic tissue. Moderate increases in vascularity were sometimes apparent. No sign of cartilage or bone erosion was observed. In contrast, the injection of xenogeneic CII induced a dense inflammatory cell infiltrate associated with increased neovascularisation and conspicuous inflammatory villus formations. None of the non-injected mice had histological abnormalities, and one out of 26 control mice injected with CFA presented a hyperplasia of the synovium in the ankle joint.

**ANALYSIS OF THE HUMORAL RESPONSE TO CII**

Mice immunised with mouse CII, bovine CII, those injected with CFA alone, or naive mice were bled at various times after the second week post-immunisation until death. The individual serum samples collected were analysed for the presence of anti-CII antibodies by ELISA. Figs 5 and 6 show the levels of antibodies directed against mouse CII obtained respectively in groups I and II. Although highly variable titres of circulating antibodies were found, all the mice injected with mouse CII developed a humoral response, whereas the serum samples from CFA injected mice or naive mice remained negative throughout the study. Interestingly, there was no relation between the levels of autoantibodies and the development of arthritis in either group of mice. Hence the females responded as well or even better than the males, irrespective of the presence of clinical signs of arthritis (Fig. 5). We observed a regular increase of antimouse CII IgG until the ninth week in group I (Fig. 5), whereas in group II the males showed a progressive decline in their antibody titres from the fifth to the fifteenth week post-immunisation (Fig. 6). It is probable that the booster intraperitoneal injection given three weeks after priming to the mice in group I, but not to the males in group II, delayed and enhanced the humoral response. Similarly, group II females that were challenged with mouse CII emulsified in IFA showed very high autoantibody levels which peaked 13 weeks after priming (Fig. 6).

All the serum samples from the mice in group I were tested for their cross reactivities with mouse and bovine CII. Several observations can be inferred from the results shown in Fig. 7. Firstly, the injection of heterologous CII elicited a humoral response that was at least 10-fold higher than that induced by homologous CII. Secondly, the antibody secreted in response to the autoantigen increased until the death of the animals at nine weeks. Thirdly, despite an important cross reactivity for CII of both origins the sera reacted best with the immunising protein in comparison with the other one. In other words, both males and females receiving mouse CII developed higher titres of antibodies reacting with mouse CII than with bovine CII, and vice versa.

**Discussion**

This paper describes the induction of an autoimmune arthritis in mice which mimics the clinical features of human RA. By use of immunisation with a purified homologous protein, thereby avoiding immune reactivity against xenogeneic determinants, we were able to analyse the disease caused by an autoantigen. The most prominent feature that distinguishes our model from the arthritis induced with heterologous CII is the fluctuation between remissions and exacerbations observed during the course of the disease. Thus in contrast with the bovine CII induced arthritis, which develops explosively one month after immunisation and never relapses after
remission, the first macroscopic signs of EAA occurred later and progressively. After an acute inflammatory episode the affected joints gradually and partially recovered, and several days or weeks later the disease recurred affecting the same or other joints. In a recent study Holmdahl et al reached identical conclusions, although their overall incidence of arthritis (30%) after administration of homologous CII to male and female mice was lower than in our experiments (58.5% in the two groups combined). Since the regimen for the induction of arthritis differed in the two studies it is possible that the immunisation protocol represents a critical point for the development of the disease. With respect to the histological features of EAA, cellular hyperplasia of the synovial membrane was always observed in the affected joints, even though the pathological changes were moderate, and the inflammatory cell infiltrate was less intense in the polyarthritis induced with the homologous than with the heterologous CII. It is possible that more severe histological lesions may be observed at the initiation or during
the course of the disease, which then regressed since, at the time of deaths animals showed only mild synovitis. This possibility is under investigation in our laboratory.

Interestingly, in our model the clinical patterns and the kinetics were highly variable from one mouse to another. Such clinical evolution resembles that of human RA, and therefore our experimental model would be a useful tool for the study of the self-perpetuating mechanisms that characterise RA. In contrast with the human disease, however, the females were significantly less susceptible to the induction of arthritis than were the males. In addition, the few females that developed arthritis after immunisation with homologous CII showed mild disease states. These findings are reminiscent of ankylosing spondylitis. The sex linkage of EAA that we observed is consistent with a recent report\textsuperscript{27} but is not supported by findings described earlier in female rats\textsuperscript{17} \textsuperscript{18} or in mice.\textsuperscript{20} It is unlikely that the weak susceptibility of females to EAA induction is related to the amount of autoantigen injected, given that a booster injection of mouse CII in IFA failed to augment the incidence of EAA, while it increased the levels of autoantibodies produced. Rather, it seems that female sex hormones exert a suppressive effect on the induction of autoimmune arthritis.\textsuperscript{28}

A critical role for antibodies against native CII in the initiation of CIA has been suggested by several investigators who demonstrated that the severity of the experimental disease was associated with a high humoral response to the immunising antigen.\textsuperscript{18} \textsuperscript{28} \textsuperscript{29} In addition, anti-CII IgG either purified from arthritic animal serum samples\textsuperscript{30} \textsuperscript{33} or monoclonal antibodies\textsuperscript{34} \textsuperscript{35} were shown to be active in transferring the disease. In the present study we found highly variable amounts of antibodies directed against homologous CII, which did not correlate with the arthritic status of the mice. Indeed, the kinetic studies performed on our two groups of mice showed that the males secreted low amounts of specific antibodies, which increased to reach a peak five to seven weeks after immunisation and then progressively declined. In contrast, both arthritic and non-arthritic females produced higher levels of anti-CII IgG than did the males. Our findings that mice developed a humoral response to CII regardless of their arthritic status argue against an antibody

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**Fig. 7** Cross reactivity of the serum samples from group 1 mice immunised with mouse CII (left side) or bovine CII (right side). Individual serum samples from males (M) and females (F) were tested for their reactivities against mouse CII (■) or bovine CII (□) six and nine weeks post-immunisation.
mediated pathogenesis when the immunising protein is derived from mouse cartilage. At present we cannot rule out the possibility that the disease is related to the secretion of particular IgG subclass autoantibodies, but is not influenced by other isotypes. Evidence suggesting that IgG2a autoantibodies are more pathogenic than other anti-CII isotypes has been reported for the arthritis induced by heterologous CII injection.31 36 This hypothesis is presently under investigation in our laboratory.

An alternative possibility regarding a potential role of type II collagen specific autoantibodies concerns the existence of different epitopes on the molecule. There is increasing evidence that the injection of pepsin solubilised CII provokes a diversity of anti-CII antibodies, some of which are directed against arthritogenic epitopes, whereas others recognise antigenic epitopes.37 38 Indeed, we, and others,27 have provided evidence that CII of xenogeneic origin induced considerably higher titres of anti-CII antibodies than did the homologous protein. This observation may reflect the existence of one or several species specific epitopes on the collagen molecule that trigger the autoimmune reactivity. Thus the susceptibility to collagen arthritis was shown to be under the control of different immune response genes according to the origin of the CII,24 37 a finding that exemplifies antigenic differences between species. The important cross reactive capacity of the antisera to bovine and mouse collagen implies that the collagen molecules from both species share immunogenic determinants. The consistently higher humoral response against the immunising protein than against the CII of the other origin supports the existence of at least one species restricted epitope.

In the light of these interpretations the failure of female mice to develop arthritis after injection of homologous CII, despite a strong humoral response, may be explained in terms of unresponsiveness to the mouse specific arthritogenic epitope(s), due either to a sex hormone linked negative control of antigen recognition28 or to an antibody mediated suppressive mechanism.39 41 Additional work is needed to elucidate this alternative hypothesis.

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