Absence of *Mycobacterium tuberculosis* DNA in synovial fluid from patients with rheumatoid arthritis

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**Abstract**

**Objectives**—To determine whether *Mycobacterium tuberculosis* DNA can be detected in synovial fluid of patients with rheumatoid arthritis (RA).

**Methods**—The polymerase chain reaction was applied to cellular components of synovial fluid.

**Results**—No evidence of *M tuberculosis* DNA was found in synovial fluid from 31 patients with RA and 13 control patients.

**Conclusion**—The findings do not support a role for persistent *M tuberculosis* infection in the pathogenesis of RA.


The aetiology of rheumatoid arthritis (RA) remains unknown, but the possible role of an infectious agent either as a trigger to the disease, or as a persistent antigenic drive to inflammatory synovitis has not been ruled out. Evidence from laboratory animals has suggested that *M tuberculosis* could be involved in the pathogenesis of arthritis. Autoimmune arthritis can be induced in MHC susceptible rats by inoculation with *M tuberculosis* and the disease can be transferred to naive animals by *M tuberculosis* specific T lymphocyte clones from inoculated rats. T cell clones isolated from Lewis rats with adjuvant-induced arthritis have been shown to proliferate in response to the 65 KDa protein of *M tuberculosis* and T lymphocytes of RA patients have been reported to show augmented reactivity to an antigen of *M tuberculosis*. IgG and IgA antibodies to the 65 KDa protein of *M tuberculosis* in RA are also reported to be significantly higher than in patients with systemic lupus erythematosus, Crohn’s disease or healthy controls, suggesting that immune responses to the 65 KDa protein may be involved in the initiation or maintenance of RA.

In the present work, we looked for the presence of *M tuberculosis* in synovial fluid of patients with RA by using the polymerase chain reaction (PCR).

**Patients and methods**

**PATIENTS**

Synovial fluid samples from 31 patients fulfilling the American Rheumatism Association Criteria (1987) for definite RA were screened by PCR, for the presence of *M tuberculosis* DNA. As a comparative group 13 patients (reactive arthritis = 4, psoriatic arthritis = 4, osteoarthritis = 2, ankylosing spondylitis = 2, gout = 1) were studied. The characteristics of the patients studied are shown in the table.

Synovial fluid was aspirated using an aseptic technique. The needle was removed and the syringe capped, coded and sent to the laboratory within two hours. The cap of the syringe containing synovial fluid was removed in a Category II, laminar flow cabinet, dedicated for this purpose. Synovial fluid was divided into aliquots of 0·5 ml and stored at –70°C.

**EXTRACTION OF DNA FROM CELLULAR COMPONENT OF SYNOVIAL FLUID**

After washing with analar water (BDH), the cell pellets from 500 μl of synovial fluid were treated with Hirt buffer (0·01 M tris, 0·01 M EDTA pH 8·0, 0·6% sodium dodecyl sulphate, SDS) and proteinase K (50 μg/ml), followed by incubation at 37°C from 12 to 18 hours. The cell lysate was then extracted three times with phenol/chloroform (1:1 mixture), and DNA was precipitated by addition of 20–50 μg of glycogen (Boehringer Mannheim), 1/10 volume of 5 M ammonium acetate and 0·8 volume of isopropanol. The DNA precipitate was washed in 70% ethanol, dried and re-dissolved in 100 μl analar water.

**POLYMERASE CHAIN REACTION USING PRIMERS FROM HUMAN APRT GENE**

The PCR reaction mixture contained 10 mM Tris pH 9, 50 mM KCl, 1·5 mM MgCl₂, 20 μM of each dNTP, 0·05 μM of each primer, 0·25 units of supertaq polymerase (HT Biotechnology Ltd) and 5 μl of template DNA.

The samples were subjected to 30 cycles of amplification in an OmniGene-Hybird Thermal Cycler as follows. Before the start of cycling the DNA was denatured for three minutes at 94°C. In each cycle denaturation was 94°C for 30 seconds, annealing of primers to the target DNA was carried at 63°C for one minute and synthesis at 72°C for one minute. At the end of 30 cycles the tubes were heated at 72°C for seven minutes.

Special care was taken to avoid contamination of samples with ampiclons. Clinical material was handled in a category II laminar flow hood, dedicated for this purpose. The preparation of PCR reaction mixture and
pipetting of samples were performed in a category II laminar flow hood situated in a separate room with restricted passage of personnel and decontamination barriers including changing of protective clothing on entering. The analysis of PCR products was performed in another laboratory room. Special emphasis was placed on preventing carryover of amplicons from post-PCR to pre-PCR facilities. To prevent aerosol contamination of pipettes aerogard tips with a barrier (Alpha Laboratories Ltd) were used in pipetting specimens.

Primers used in this study were designed by using PCR and Oligo Primer Design Program (Scientific Educational Software, USA). The DNA sequences of primers employed were as follows (reading from 5' to 3').

Human APRT gene (amplifies: 329 bp from bp 1198–1527)

(1) CTTCCTGTTCTCTCTCGAG
(2) TGGCCTGAAGCTGTGATGTT

POLYMERASE CHAIN REACTION USING PRIMERS FROM A REPETITIVE DNA SEQUENCE SPECIFIC FOR M TUBERCULOSIS

Each DNA sample from cellular components of synovial fluid was spiked with 5 pg of M tuberculosis DNA to investigate the presence of PCR inhibitors in it. PCR was performed on spiked and unspiked samples as described by Eisenach et al.6

DETECTION OF AMPLIFIED PRODUCTS

Fifteen μl of each PCR product was analysed by gel electrophoresis in a 2% agarose gel using tris borate buffer (89 mM tris, 89 mM boric acid pH 7.8). The amplified products were visualised by staining the gel with ethidium bromide (0·5 μg/ml in tris borate buffer) and inspected under UV illumination.

Results

SUITABILITY OF DNA FOR AMPLIFICATION

To exclude possible false negative results caused by absences of template DNA, PCR was performed using primers specific for the human adenosine phosphoribosyl transferase (APRT) gene. Of the 44 synovial fluid samples investigated in this study, 37 contained amplifiable DNA. An example of the results is shown in fig 1.

PCR TO DETECT M TUBERCULOSIS DNA IN SYNOVIAL FLUID

Spiked samples

PCR was performed on all 44 samples spiked with M tuberculosis DNA, using primers specific for M tuberculosis under highly stringent PCR conditions. The sensitivity and specificity of PCR assay used in this study has been described elsewhere.6 All spiked samples gave positive results. An example of the results is shown in fig 2.

Unspiked samples

PCR was also performed on all 44 unspiked samples. None of the unspiked synovial fluid sample contained M tuberculosis DNA. An example of the results is shown in fig 3.
absence of mycobacterium tuberculosis DNA in synovial fluid from patients with rheumatoid arthritis

To ensure the reproducibility and reliability of PCR, positive and negative controls were included in each experiment. Positive control for PCR using primers from human APRT gene was DNA extracted from a pelvic lymph node and for PCR using primers from repetitive DNA sequence of M. tuberculosis was pure growth of recent clinical isolate of M. tuberculosis. Negative controls contained all the ingredients of PCR except DNA which was replaced by equal volume of anular water. All controls gave appropriate results (figs 1, 2 and 3).

Discussion

The potential role of microbes in the aetiology of RA has been highlighted by the demonstration of microbial components at the site of inflammation in reactive arthritis. This has been documented in arthropitides following infection with Chlamydia, Yersinina, Salmonella and Shigella.11-10 Borrelia burgdorferi DNA has also been detected in synovial fluids of patients with Lyme arthritis by PCR.11

Although evidence from laboratory animals has suggested a possible role of M. tuberculosis in RA, in this study its DNA was not detected by PCR in the cellular components of any synovial fluid from 31 RA patients or 13 cases with other forms of arthritis. Isolation of cytotoxic T cells clones crossreactive with M. bovis and human hsp 60 from the joints of RA patients suggests a role of autoreactive T cells directed against self-hsp in the pathogenesis of some forms of chronic inflammatory arthritis.12

However, this view has been questioned. First, it appears that synovial fluid T cells reactive with hsp 60 in human RA generally recognise nonconserved bacterial rather than shared (or human) epitopes;13 second, at least some T cells responses to recombinant mycobacteric hsp 60 may be directed against contaminating E.coli antigens rather than hsp 60 itself;14 finally the frequency of T cells reactive with mycobacterial hsp 60 appears to be virtually identical in blood and synovial fluid of patients with RA casting doubt on previous suggestions of local expansion of hsp-reactive T cells in the disease.15 The role of molecular mimicry between M. tuberculosis and humans hsp 60 proteins is far from clear.

The PCR assay used in this study is highly sensitive and specific. It is able to detect a single copy of M. tuberculosis chromosome from a mixture of human and bacterial DNA.4 Extreme care was taken to code the samples, ensuring that the reactions were carried out without bias, with the inclusion of positive and negative controls and comparison of results between different PCR runs. While false positive results are a common source of concern in PCR, a frequent source of false negative results in survey of this type is sampling variability, which results in either no DNA or DNA which is unsuitable for the PCR reaction. This possibility was ruled out by employing primers from a cellular gene (APRT). Thirty seven of forty four synovial fluid samples gave positive results, showing that the DNA from these samples was sufficient to act as a PCR template. The possibility of PCR inhibitors in DNA prepared from synovial fluid was also investigated by spiking each sample with M. tuberculosis DNA. All 44 samples including seven samples which gave negative result with PCR using primers from APRT gene, produced an expected band of 123 bp showing absence of inhibitors in DNA preparation (fig 2). This shows that the DNA extraction method used in this study removes PCR inhibitors and does not reduce the amplifiability of the DNA template. Seven samples which gave negative result with PCR using primers from APRT gene shows the absence of DNA rather than the presence of PCR inhibitors in amplification reaction. Therefore a negative result in the rest of 37 samples, using the primers for M. tuberculosis, would probably be due to absence of M. tuberculosis DNA and not to a total lack of DNA.

Persistent infection at sites other than joints can cause reactive arthritis.16 This possibility has not been excluded by this study. Absence of M. tuberculosis DNA in synovial fluid from patients with RA also does not exclude the presence of M. tuberculosis DNA in synovial membranes. Although the techniques used in this study could be used to identify the presence of microbial DNA in synovium, it was not considered ethical to collect samples of synovium from the patients included in this study.

Our findings do not support the persistent involvement of M. tuberculosis in the aetio-pathogenesis of RA. However, its role as a triggering factor in a genetically or otherwise predisposed host cannot be ruled out.

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