The value of synovial fluid analysis in the assessment of knee joint destruction in arthritis in a three year follow up study

S Åman, J Risteli, R Luukkainen, L Risteli, M Kauppi, P Nieminen, M Hakala

Abstract

Objectives—To assess the predictive significance of synovial fluid (SF) analysis for progressive radiological knee joint destruction in arthritis.

Methods—Altogether 55 patients with arthritis and knee joint effusion were included in the study. The diagnosis was rheumatoid arthritis (RA) for 44 of them, chronic seronegative spondylarthropathy for seven and juvenile rheumatoid arthritis for four. The mean age of the patients was 51.8 (SD 14.9, range 19–82) years, and the mean duration of disease 10.9 (SD 9.2, range 0.5–37) years. In addition to the routine laboratory tests, different markers of collagen synthesis and breakdown in serum and SF were assessed. The radiological grade of the knee joint was assessed by Larsen’s method at the baseline and after a three year follow up.

Results—During the follow up, Larsen’s grade deteriorated in 22 (40%) patients. These patients had a significantly higher median level of cross linked carboxyterminal telopeptide of type I collagen (ICTP) in SF at entry than those who had a stable index (p = 0.035). Serum ICTP did not have any predictive value for a specific joint. The median levels of total SF leucocytes (p = 0.012) and the subgroup of polymorphonuclear leucocytes (p = 0.018) were higher in the patients with a stable Larsen’s index. However, the relation of SF leucocyte level to radiological progression could not be confirmed in the RA group.

Conclusion—It is concluded that SF analysis may help in the identification of patients with inflammatory arthritis who are at risk for progressive destruction in a particular joint. A high total SF leucocyte level is not necessarily associated with a poor prognosis. Instead, a high SF ICTP level seems to reflect accelerated bone degradation.

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There is increasing evidence to suggest that synovial fluid (SF) analysis may help in analysing the further destruction of an individual joint in patients with arthritis.1–4 Much interest has been shown in the measurement of the synthesis and breakdown of different components of bone and cartilage.1–5 Saxne et al studied a group of patients with rheumatoid arthritis (RA) from whom knee joint SF had been aspirated 10 years previously and found a correlation between the SF proteoglycan concentrations and the degree of subsequent radiographic joint destruction.1 In two other studies, an increasing SF bone sialoprotein (BSP) concentration was accompanied by increasing degrees of knee joint damage.1,2 BSP is a bone specific macromolecule that constitutes some 12% of the non-collagenous proteins in bone, being hence a major component of this group of proteins.8 In a longitudinal study by Månsson et al, the concentrations of SF aggrecan—that is, the cartilage oligomeric matrix protein (a non-collagenous matrix protein)—were initially highest in the group of RA patients with developing joint destruction.9

We have earlier measured the breakdown of type I collagen using an assay of cross linked carboxyterminal telopeptide (ICTP)10 in the SF of 59 unselected patients with RA and hydropsy in a knee joint. SF ICTP was clearly related to the radiographic findings in the corresponding knee joint, as the patients with gross bone deformation had the highest SF ICTP concentrations. In addition, the markers for type I and type III collagen synthesis, the aminoterminal propeptides of type I (PINP)11 and type III procollagen (PIIINP)12, when measured from SF, correlated clearly with each other and with SF ICTP.13 In the present series, we looked for a connection between SF analysis at the baseline and radiological status after a three year follow up in 55 patients with arthritis.
and interassay coefficients of variation of the three markers tested with human serum samples are less than 10%.\textsuperscript{10–12} An aliquot of each serum sample was centrifuged within one hour after aspiration at 1800 g for two periods of 20 minutes. Intra-assay variation of the SF samples was 4.4% for PINP and 5.7% for PII-INP. The SF samples of the series were not centrifuged. We later tested the effect of centrifugation on the results in five additional patients. Centrifugation was found to decrease the levels of the markers of collagen metabolism in SF to the same extent as does centrifugation of plasma into serum. With the exception of one sample each, the level of the markers after centrifugation was lower compared with native samples by amounts varying from 6.3 to 11.7% for SF ICTP and from 0.2 to 1.4% for SF PII-INP.

### Statistical analysis

The data were analysed on a personal computer using SOLO statistical software.\textsuperscript{16 17} Mann-Whitney test, Kruskal-Wallis test and \( \chi^2 \) statistics were used as appropriate.

### Results

At entry, SF ICTP correlated positively with Larsen’s index. Table 1 shows the distribution of Larsen’s grades at entry and at the three year visit. The median SF ICTP concentrations in the baseline Larsen’s grade groups 0–4 were 10.8, 15.5, 12.5, 18.3 and 33.8 \( \mu \)g/l, respectively (\( p = 0.009 \)). The radiological index deteriorated in 22 (40%) patients, but remained stable in 23 (42%). Table 2 shows the median baseline values of different markers in both groups. The median SF ICTP concentration was higher in the patients with a deteriorating Larsen’s grade than in those with a stable one (\( p = 0.035 \)). When tested only in the patients with RA (\( n = 44 \)), an equal difference in SF ICTP was found between the above groups—that is, the median concentration of SF ICTP was 14.0 \( \mu \)g/l (range 2.8–42.3 \( \mu \)g/l) for the 25 RA patients with stable disease and 18.3 \( \mu \)g/l (range 7.1–64.6 \( \mu \)g/l) for the 19 subjects whose disease was deteriorating, though the difference did not reach statistical significance (\( p = 0.063 \)). Instead, both the median levels of SF total leucocytes (\( p = 0.012 \)) and polymorphonuclear leucocytes (\( p = 0.018 \)) were statistically higher in the patients with a stable grade. A comparison of the diagnostic subgroups showed that the association did not hold in the RA group, but was more evident in the patients with stable disease.
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For stable disease 11.0 × 10^9/l for progressive disease in the RA group, while the corresponding values in the other diagnostic groups were 12.6 × 10^9/l and 2.2 × 10^9/l. There was no statistically significant difference for SICTP, SP IIINP, SF PINP or SF PIIINP between the above groups either in the whole series or in the diagnostic subgroups. The mean SF:S ratio of ICTP was 3.1 (SD 1.8) for the patients with a stable Larsen's grade, and 4.5 (3.3) for the patients with a deteriorating Larsen's grade (NS).

There was no statistical difference between the patients with a stable or deteriorating Larsen's grade with regard to treatment with DMARDs or peroral corticosteroid therapy during the follow up.

During the follow up, 13 (42 %) of the 31 patients whose Larsen's grade remained stable and 15 (71%) of the 21 whose grade deteriorated needed one or more corticosteroid injections (including chemical synovectomy in five cases) (p = 0.036); for three patients the follow up data were missing.

Discussion

Markers of collagen synthesis and degradation measured in serum and SF reflect the grade of synovitis and tissue breakdown in RA. The earlier studies to measure collagen degradation were based on the use of urine samples including assays of the amino acid 4-hydroxyproline, pyridinoline and deoxypyridinoline cross links, and showed controversial results as to the association with radiological progression in hands and feet.

In the present series, we used serum radioimmunoassays to measure the synthesis and degradation of type I collagen (PINP and ICTP) and the synthesis of type III collagen (PIINP) in patients with different forms of inflammatory arthritis. Type I collagen is the most abundant protein species in the human body. Most of it is present in bones, where it accounts for about 90% of the organic matrix, and it is also the major matrix protein in tendons, ligaments, and soft connective tissues. Type III collagen is the second most abundant collagen type and is found in soft connective tissue.

The only study to compare ICTP with the urinary assays of type I collagen degradation was a cross sectional analysis of five different markers of collagen degradation in RA, where the serum ICTP and urinary pyridinoline (PYD) levels were found to be superior to the other markers in discriminating between the RA and control groups. PYD cross links arise not only from type I collagen degradation, but also from type II collagen degradation in articular cartilage.

At the beginning of the present follow up, the mean SF concentrations of the collagen derived substances studied were high in our patients, and their concentrations correlated with the grade of destruction in the joints.

During the three year follow up, Larsen's grade deteriorated in 22 (40%) patients. These patients had a significantly higher SF ICTP level at entry than the ones who had a stable grade. Serum ICTP did not have the same predictive significance, nor did the serum or SF assays for PINP and PIINP. Serum ICTP reflects the type I collagen metabolism in the whole body, but in our earlier community-based RA series, 26% of the patients with initially increased serum ICTP values required total joint replacement surgery of at least one joint during a three year follow up, compared with 2% of the patients with initially normal ones. However, when estimating the outcome of one particular joint, such as a knee in this study, it seems more logical to measure ICTP from SF than from serum.

Interestingly, the total number of SF leucocytes and the number of polymorphonuclear leucocytes were here higher in the patients with a stable erosive grade than in those with a progressive one (p = 0.012 and p = 0.018, respectively). A comparison of the diagnostic subgroups showed that the association did not hold true in the RA group, but was more pronounced in the other inflammatory arthropathies. The non-RA patient group represented different arthritides and included a small number of patients (n = 11), which decreases reliability of the finding. However, it is to be noted that a recent study on the relation of synovial biopsy findings to SF analysis in 33 patients with inflammatory arthritis suggested that SF cell counts reflect the activity of acute synovial inflammation but not that of chronic inflammation.

In the present series, baseline SF ICTP correlated positively with Larsen's grade, and those with the highest SF ICTP levels most often deteriorated during the follow up. It should be noted that those with a deteriorating Larsen's grade were more often treated with intra-articular corticosteroids than those with a stable grade (p < 0.05). Thus, once the breakdown process has started, it is seldom stopped by the present treatment. The role of local corticosteroids in accelerating joint destruction can only be speculated.

We conclude that SF analysis may help in assessing the disease process of a particular joint in patients with inflammatory arthritis. According to our results a high SF ICTP level seems to reflect accelerated synovial tissue and bone degradation. Instead, our preliminary results indicate that a high SF leucocyte level is not necessarily associated with a poor prognosis. More studies are needed to clarify the question of the relation between the level of SF leucocytes and the development of further joint destruction in patients with inflammatory arthritis.


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