'Dialysis related arthropathy': a survey of 95 patients receiving chronic haemodialysis with special reference to $\beta_2$ microglobulin related amyloidosis

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SUMMARY Ninety five patients receiving chronic haemodialysis (CHD) were surveyed to determine the prevalence of rheumatic disease and, where possible, its aetiology. At least three distinct rheumatic syndromes were identified—a group of patients with a syndrome consisting of large and medium joint synovial swelling, restricted hips and shoulders, tenosynovitis, carpal tunnel syndrome, and bone cysts due to deposition of $\beta_2$ microglobulin related amyloid (AM$_{\beta_2m}$); a second group with erosive azotaemic osteoarthropathy; and a third group with age related degenerative disease of small, large, and axial joints. The data presented suggest that in patients receiving CHD (a) the prevalence of AM$_{\beta_2m}$ deposition and the associated syndrome increases with duration of dialysis, but in patients who have been dialysed for more than 10 years the risk of developing AM$_{\beta_2m}$ is related to age; (b) AM$_{\beta_2m}$ deposition in subchondral cysts, but not synovium, causes joint destruction; also, AM$_{\beta_2m}$ may be more prone to deposition in synovium of joints already damaged by other processes; (c) in the absence of synovial iron deposition synovial AM$_{\beta_2m}$ is not associated with an inflammatory infiltrate; (d) hyperparathyroidism and perhaps other factors such as synovial iron deposition are probably more important than AM$_{\beta_2m}$ as causes of peripheral joint degeneration and destructive spondyloarthropathy in patients receiving CHD.

A number of skeletal and articular abnormalities have been described in patients undergoing chronic haemodialysis (CHD). These include renal osteodystrophy, avascular necrosis of bone, crystal induced arthritis, periarticular calcification,1-3 and recurrent haemarthrosis of shoulders.2 3 A syndrome, comprising carpal tunnel syndrome (CTS),3-8 chronic large joint arthropathy,3 9 and bone cysts,10 has also been recognised as a major complication of CHD. Recent studies have demonstrated the deposition of amyloid in the carpal tunnel, bone cysts, synovium, and tendons of such patients.8 10-14 The major component of the amyloid has been identified as $\beta_2$ microglobulin,15-25 but amyloid P component is also present.21 A suggested designation is ‘AM$_{\beta_2m}$’.19 AM$_{\beta_2m}$ has also been identified in synovial fluid24 and vertebral discs.25 AM$_{\beta_2m}$ deposits have been infrequently described in other major organs12 23 and, where present, are usually small and perivascular.26 27 A similar tropism for synovial tissue has been described previously for AL-type amyloid,28 but AA-type amyloid usually spares synovium and bone. A minor degree of synovial amyloidosis occurs with increasing age.29

Some bone deposits of AM$_{\beta_2m}$ have been responsible for fractures,30 31 and it has been suggested that AM$_{\beta_2m}$ is responsible for the erosive arthropathy11 13 21 23 and spondyloarthopathy25 which may occur in patients undergoing CHD. It is not certain, however, whether amyloid alone or other factors such as synovial iron deposition contribute to the large and medium joint arthropathy associated with this syndrome.3 32

The present study was undertaken to assess the prevalence of rheumatic disease and, where possible, its relation with tissue deposition of AM$_{\beta_2m}$ or other factors in a defined population of patients receiving CHD.

Patients and methods

The patient population consisted of all 118 patients accepted for publication 27 July 1988.

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receiving CHD under the care of the Queen Elizabeth Hospital renal unit during a defined nine month period (May 1986–January 1987). Of these, 23 patients were excluded from the study: seven had received renal transplants, two died before being reviewed, and a further 14 patients who were dialysed at remote regional centres (>250 miles) were not seen for logistic reasons. Ninety five patients (40 female, 55 male) were available for review.

All patients were reviewed for symptoms of musculoskeletal disease or symptoms related to systemic amyloidosis using a structured questionnaire and then underwent a complete physical examination performed by one of two rheumatologists (NH, R van den B). The presence of joint ‘restriction’ was defined as loss of 30% or more of the normal range of movement and ‘synovitis’ as the presence of palpable synovial thickening or clinically detectable effusion, or both. CTS was or had been diagnosed on the basis of a typical clinical history, physical findings, and nerve conduction studies. In all cases details of dialysis, cause of renal failure, and parathyroid status were obtained by review of case records.

All patients had radiographs taken of hands, wrists, shoulders, hips, knees, feet, and spine. These were reviewed by two radiologists (LA, MA) experienced in reporting renal bone disease. A radiological diagnosis of renal osteodystrophy was made if any of the following were present: subperiosteal resorption, tuft resorption, distal clavicular erosions, rugger jersey spine or other typical bony sclerosis, or coarse bony trabeculation. A diagnosis of significant active hyperparathyroidism was based on finding either (a) typical radiological changes and parathyroid hormone >1000 pmol/l or (b) a bone biopsy specimen showing active bone resorption. Nodal erosive osteoarthrosis of hands was defined as loss of joint space, osteophyte formation, and erosive bone damage, usually affecting interphalangeal or first carpometacarpal joints. Erosive azotaemic osteoarthropathy was defined as the presence of marginal joint erosions, usually affecting metacarpophalangeal or interphalangeal joints, and occasionally larger joints, in the absence of joint space narrowing.

The following biochemical studies were performed on serum or plasma from all patients: serum \(\beta_2\) microglobulin before and after dialysis, C reactive protein, ferritin, serum electrophoresis, rheumatoid factor, and antinuclear factor. Parathyroid hormone was measured by immunoassay for the C terminal fragment.

Iliac crest bone biopsy results were reviewed from 68/95 patients. All biopsies had been carried out within 18 months of the study and most within 12 months. The remaining patients had either declined to undergo bone biopsy or the available biopsy had been performed more than 18 months previously. Other histological specimens were available from 12 patients and synovial fluid from a further seven patients. These were examined for the presence of amyloid by Congo red staining. Where Congophilia was identified the presence of \(\beta_2\) microglobulin was sought using rabbit antihuman \(\beta_2\) microglobulin antibody and the peroxidase-antiperoxidase technique (Dakopatt). Tissues containing either AA or AL-type amyloid were used as negative controls for immunoperoxidase studies. Deposition of iron in tissues was sought using Perls’ stain. Synovial fluid was centrifuged and the sediment examined for the presence of crystals by polarising microscopy, or deposits of \(\text{AM}_{\beta_2}\) using the immunoperoxidase technique followed by Congo red.

Results

Patients

Patients were divided into three cohorts according to duration of haemodialysis: group A = 0 to 4.9 years (n=49), group B = 5 to 9.9 years (n=32), and group C = 10 years (n=14). The mean (SD) ages were: group A 55.8 (11.9), group B 46.8 (13.3), group C 52.1 (13.2) years.

The causes of renal failure included AA-type amyloid in one patient and multiple myeloma in another. They had received dialysis for two years and one year respectively and neither had arthropathy, bone cysts, or CTS.

Patients had been treated mostly with cuprophane membrane dialysers. Reverse osmosis has been used for the last 10 years ensuring dialysate aluminium concentrations <10 \(\mu\)mol/l.

Clinical and radiological findings

Joint symptoms

Pain and stiffness affecting peripheral and axial joints were very common and the frequency rose with increasing duration of dialysis (Table 1).

Large and medium size joints

The frequency of joint restriction, synovitis, and effusions in large and medium size joints rose with duration of dialysis and affected over 60% of the patients in group C (Tables 1 and 2).

Synovitis was most common in the knee (Table 2); in five of these patients there was no radiological change, in five osteoarthritis, in one a large subchondral tibial cyst later shown to contain \(\text{AM}_{\beta_2}\) (see below), and in one loss of articular cartilage. In
one patient with an acute painless effusion haemarthrosis was found but no organisms or crystals. Restriction of knee movements was an infrequent finding (Table 2).

Chronic synovitis of shoulders was less common than restriction (Table 2), and these signs occurred together in two patients. In two patients with acute painful exacerbation of chronic synovitis haemarthrosis was found; from one of these Escherichia coli was isolated but in the other no explanation was found for bleeding. Several patients had radiological evidence of major rotator cuff attrition, often associated with loss of articular cartilage and secondary osteoarthrosis, but only one had synovitis. The frequency of significant rotator cuff disease increased with age (Table 3) but not with duration of dialysis (Table 4).

Restriction of hip movement was seen in 17 patients. This was associated with no radiological abnormality in 10, femoral head cysts in two, osteoarthritis in three and avascular necrosis with secondary osteoarthrosis in three patients, of whom two had required total hip replacement. Asymptomatic femoral neck cysts or avascular necrosis respectively were seen in two and three patients each. Thus no radiological abnormality was identi-

### Table 1 Clinical abnormalities. The number (%) of patients affected in each cohort* is shown

<table>
<thead>
<tr>
<th>Joint/Abnormality</th>
<th>A (n=49)</th>
<th>B (n=32)</th>
<th>C (n=14)</th>
<th>p Value†</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Large/medium joints</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain/stiffness</td>
<td>14 (29)</td>
<td>12 (38)</td>
<td>11 (79)</td>
<td>&lt;0-001</td>
</tr>
<tr>
<td>Synovitis/effusion</td>
<td>2 (4)</td>
<td>5 (16)</td>
<td>8 (57)</td>
<td>&lt;0-001</td>
</tr>
<tr>
<td>Restriction</td>
<td>11 (22)</td>
<td>12 (38)</td>
<td>9 (64)</td>
<td>&lt;0-01</td>
</tr>
<tr>
<td><strong>Small joints (hands)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain/stiffness</td>
<td>26 (53)</td>
<td>20 (63)</td>
<td>13 (93)</td>
<td>&lt;0-01</td>
</tr>
<tr>
<td>Synovitis/effusion</td>
<td>0 (0)</td>
<td>1 (3)</td>
<td>1 (7)</td>
<td>NS</td>
</tr>
<tr>
<td>Osteoarthritis</td>
<td>12 (24)</td>
<td>4 (13)</td>
<td>3 (21)</td>
<td>&lt;0-01</td>
</tr>
<tr>
<td>Flexor tenosynovitis</td>
<td>1 (2)</td>
<td>0 (0)</td>
<td>3 (21)</td>
<td>&lt;0-01</td>
</tr>
<tr>
<td><strong>Axial joints</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain/stiffness</td>
<td>13 (27)</td>
<td>12 (38)</td>
<td>6 (43)</td>
<td>NS</td>
</tr>
<tr>
<td>Restriction</td>
<td>6 (12)</td>
<td>6 (19)</td>
<td>4 (29)</td>
<td>NS</td>
</tr>
<tr>
<td>Carpal tunnel syndrome</td>
<td>0 (0)</td>
<td>1 (3)</td>
<td>6 (43)</td>
<td>&lt;0-001</td>
</tr>
</tbody>
</table>

*Duration of dialysis for each group was as follows: A=0-4-9 years; B=5-9-9 years; C=>10 years.
†x² test, group C v group A.

### Table 2 Distribution of synovitis and restriction in peripheral joints (number of patients affected)

<table>
<thead>
<tr>
<th>Joint</th>
<th>Synovitis</th>
<th>Restriction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Knees</td>
<td>12</td>
<td>4</td>
</tr>
<tr>
<td>Shoulders</td>
<td>3</td>
<td>14</td>
</tr>
<tr>
<td>Elbows</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>Wrists</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Hips</td>
<td>—</td>
<td>17</td>
</tr>
<tr>
<td>Subtalar</td>
<td>—</td>
<td>6</td>
</tr>
</tbody>
</table>

Total number of affected patients 15 32

### Table 3 Relation between radiological changes of osteoarthritis and age. The number (%) of patients affected is shown

<table>
<thead>
<tr>
<th>Age (years)</th>
<th>Number</th>
<th>Large joint OA*</th>
<th>Cartilage erosion without OA</th>
<th>Erosive nodal OA</th>
<th>Major rotator cuff disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>20-39</td>
<td>24</td>
<td>1 (4)</td>
<td>0 (0)</td>
<td>2 (8)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>40-59</td>
<td>39</td>
<td>12 (21)</td>
<td>2 (5)</td>
<td>5 (13)</td>
<td>2 (5)</td>
</tr>
<tr>
<td>60-79</td>
<td>32</td>
<td>10 (31)</td>
<td>4 (13)</td>
<td>7 (22)</td>
<td>7 (22)</td>
</tr>
</tbody>
</table>

*OA=osteoarthritis.

### Table 4 Number (%) of patients with radiological abnormalities, analysed by cohorts*

<table>
<thead>
<tr>
<th>Abnormality</th>
<th>A (n=49)</th>
<th>B (n=32)</th>
<th>C (n=14)</th>
<th>p Value†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bone cysts</td>
<td></td>
<td></td>
<td></td>
<td>&lt;0-001</td>
</tr>
<tr>
<td>In any site</td>
<td>5 (10)</td>
<td>9 (28)</td>
<td>7 (50)</td>
<td>&lt;0-001</td>
</tr>
<tr>
<td>In carpi</td>
<td>4 (8)</td>
<td>7 (22)</td>
<td>4 (29)</td>
<td>&lt;0-05</td>
</tr>
<tr>
<td>Renal osteodystrophy</td>
<td>12 (24)</td>
<td>9 (28)</td>
<td>9 (64)</td>
<td>&lt;0-001</td>
</tr>
<tr>
<td>Erosive azotaemic osteoarthropathy</td>
<td>6 (12)</td>
<td>4 (13)</td>
<td>4 (29)</td>
<td>NS</td>
</tr>
<tr>
<td>Small joints of hands</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OA≠total</td>
<td>15 (31)</td>
<td>12 (38)</td>
<td>4 (29)</td>
<td>NS</td>
</tr>
<tr>
<td>OA-erosive</td>
<td>7 (14)</td>
<td>6 (19)</td>
<td>1 (7)</td>
<td>NS</td>
</tr>
<tr>
<td>OA-non-erosive</td>
<td>8 (16)</td>
<td>6 (19)</td>
<td>3 (21)</td>
<td>NS</td>
</tr>
<tr>
<td>Large/medium joints</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OA (excludes AVN)</td>
<td>13 (27)</td>
<td>7 (22)</td>
<td>5 (36)</td>
<td>NS</td>
</tr>
<tr>
<td>AVN</td>
<td>1 (2)</td>
<td>5 (16)</td>
<td>0 (0)</td>
<td>NS</td>
</tr>
<tr>
<td>Cartilage erosion without OA</td>
<td>5 (10)</td>
<td>0 (0)</td>
<td>1 (7)</td>
<td>NS</td>
</tr>
<tr>
<td>Axial joints</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spondylosis or apophyseal joint OA</td>
<td>3 (6)</td>
<td>3 (9)</td>
<td>4 (29)</td>
<td>&lt;0-02</td>
</tr>
<tr>
<td>Rotator cuff disease§</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Major</td>
<td>5 (10)</td>
<td>3 (9)</td>
<td>1 (7)</td>
<td>NS</td>
</tr>
<tr>
<td>Minor</td>
<td>4 (8)</td>
<td>5 (16)</td>
<td>2 (14)</td>
<td>NS</td>
</tr>
</tbody>
</table>

*Duration of dialysis for each group was as follows: A=0-4-9 years; B=5-9-9 years; C=>10 years.
†x² test, group C v group A.
§OA=osteoarthritis; AVN=avascular necrosis.
§Major=upward subluxation of humeral head; minor=rotator cuff calcification or sclerosis of greater tuberosity.
fied in over half of the patients (59%) with hip restriction.

The prevalence of large and medium joint osteoarthritis (excluding cases of avascular necrosis), or loss of articular cartilage without osteoarthritis, increased with age (Table 3) but not duration of dialysis (Table 4). Two of the six patients with avascular necrosis had not received corticosteroids.

Small joints

Small joint symptoms were frequent and increased with duration of dialysis (Table 1). Symptoms were associated with osteoarthritis in 31 patients, erosive azotaemic osteoarthropathy in six patients, metacarpophalangeal joint synovitis in two, and flexor tenosynovitis in four, but no cause was identified in 16.

In those with erosive osteoarthritis pain, instability, and deformity of the interphalangeal or first carpometacarpal joints often caused significant functional disability. Restriction of finger movements was not usually a significant functional problem except in those with flexor tenosynovitis (see below). The frequency of objective clinical or radiological signs of osteoarthritis increased with age but not with increasing duration of dialysis (Tables 1, 3, and 4). Two patients had developed erosive osteoarthritis at a young age (less than 30 years), suggesting that factors associated with renal failure had contributed. Neither had any occupational or other physical risk factor which might have contributed.

Fourteen patients had erosive azotaemic osteoarthropathy, of whom two also had osteoarthritis. In patients with erosive azotaemic osteoarthropathy small erosions were seen adjacent to the metacarpophalangeal and proximal interphalangeal joints, but in two cases were also noted adjacent to the tibial plateau and glenoid fossa respectively. Only six patients with erosive azotaemic osteoarthropathy had small joint symptoms. There was no significant correlation either between the presence of osteoarthritis and erosive azotaemic osteoarthropathy (χ² test; p>0.5) or between small joint symptoms and erosive azotaemic osteoarthropathy (χ² test; p>0.5). The possible relation between erosive azotaemic osteoarthropathy and renal osteodystrophy is discussed below.

A further two patients with small joint symptoms had metacarpophalangeal joint synovitis, of whom one had ulnar deviation. Both had negative tests for rheumatoid factor and antinuclear factor and normal plasma protein electrophoresis, but one had erosive azotaemic osteoarthropathy. A further four patients had flexor tenosynovitis of the fingers associated with marked tendon crepitus and im-

Axial joints

Restriction of spinal movement was found in 16 patients and was often sufficiently severe as to suggest the possibility of ankylosing spondylitis. The frequency of symptoms and signs rose with duration of dialysis. In six patients the stiffness affected the whole spine, in five it affected the cervical spine alone, in three the thoracolumbar spine, and in two the lumbar spine alone. In one of these patients lumbar disc protrusion and, subsequently, synovial cysts arising from lumbar apophyseal joints caused lumbar root compression requiring operative intervention. The synovium of these joints, and to a lesser extent the degenerating disc, was found to be densely infiltrated with AM₉₂m. Further necropsy findings are described below. In three patients with, and seven without, clinical signs there was radiological evidence of degenerative disc or facet joint disease, and none had evidence of ankylosing spondylitis. Two of these patients had ‘destructive’ disc lesions with loss of adjacent end plate, which resembled discitis. These occurred at T4/5, L3/4, and L4/5 in one, and at L4/5 in the other.

Carpal tunnel syndrome

CTS was diagnosed in seven patients, of whom six had undergone dialysis for over 10 years (Table 1); several had had repeated operations for carpal tunnel decompression. All seven patients with CTS had large or medium joint arthropathy, which was associated with flexor tenosynovitis in two and small joint synovitis in one. All had bone cysts.

Bone cysts

Bone cysts were seen with increasing frequency with duration of dialysis, and 50% of patients in group C were affected (Table 4). These were usually in the

<table>
<thead>
<tr>
<th>Affected bones</th>
<th>Number of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carpi</td>
<td>12</td>
</tr>
<tr>
<td>Tibia</td>
<td>1</td>
</tr>
<tr>
<td>Femur (head or neck)</td>
<td>4</td>
</tr>
<tr>
<td>Humerus (head)</td>
<td>1</td>
</tr>
<tr>
<td>Radius (distal)</td>
<td>3</td>
</tr>
<tr>
<td>Navicular</td>
<td>1</td>
</tr>
<tr>
<td>Phalanges</td>
<td>4</td>
</tr>
<tr>
<td>Total number of affected patients</td>
<td>21</td>
</tr>
</tbody>
</table>

Table 5 Distribution of bone cysts (number of patients affected)
carpi (Table 5) (Fig. 1a), but long bones were also involved; one cyst affecting the tibial plateau required curettage and bone grafting to prevent fracture (Fig. 1b). Serial radiographs of a cyst in the femoral head were also obtained (Fig. 1c). In another, a large subchondral cyst in the head of the third proximal phalanx was associated with degenerative change in the adjacent joint. As noted above, there was a close association between the presence of bone cysts and CTS.

Renal osteodystrophy
Twenty nine patients had active hyperparathyroid-
The prevalence of typical radiographic duration of renal osteodystrophy were 25 undergone parathyroidectomy and no longer had active disease. Of the 68 available biopsy specimens, 25 were normal or had very mild abnormalities, 11 showed osteomalacia. 24 active hyperparathyroidism, and eight showed evidence of previous hyperparathyroidism. Seven patients had small deposits of aluminium, and three of these had osteomalacia.

There was no statistical correlation between the presence of active hyperparathyroidism and bone cysts, erosive azotaemic osteoarthropathy, osteoarthritis or loss of articular cartilage in large or medium joints, or both, or erosive nodal osteoarthritis: 9/21 patients with bone cysts, 3/14 with erosive azotaemic osteoarthropathy, 12/30 with large or medium joint osteoarthritis or cartilage loss, and 8/14 with erosive nodal osteoarthritis respectively had hyperparathyroidism ($\chi^2$ test; $p>0.1$). Both the patients with destructive intervertebral disc lesions had rugger jersey spines, and one had 'active' hyperparathyroidism.

HISTOLOGY

One or more histological or synovial fluid specimens were available in life from a total of 15 patients. The indications for obtaining these were carpal tunnel release (seven patients), excision of bone cyst (one), and excision of iliopsoas bursa (one). Synovial fluid aspiration was performed in three of these patients and in a further seven patients with chronic large joint synovitis. Necropsy was performed on six patients, including two of the above 15 patients. Tissues in which AM$_{\beta_{2m}}$ deposition was identified by Congo red staining and by immunoperoxidase for $\beta_2$ microglobulin included carpal tunnel structures, synovium, and bone (Table 6).

**Histology and necropsy results**

In six of the seven patients with CTS AM$_{\beta_{2m}}$ was present in various carpal tunnel structures, including the flexor retinaculum, flexor tendon, and sheath. In the seventh patient dense fibrous tissue was seen in the retinaculum.

Electron microscopy studies on material from bone cysts and synovium demonstrated curvilinear fibrils (Fig. 2).

Necropsy material was available from six patients. One patient, who had received dialysis for 14 years, had severe dialysis associated arthropathy with bone cysts, recurrent CTS, synovitis of wrists and knees, restriction of shoulders, wrists, hips, and spine, and rotator cuff attrition. As mentioned above, apophysal joint swelling and disc degeneration had caused lumbar nerve root compression. At nec-

Fig. 2 Electron microscopy of amyloid identified by Congo red staining in the synovium of a knee. Short curvilinear fibrils of amyloid are seen.

### Table 6  Sites of AM$_{\beta_{2m}}$ deposition

<table>
<thead>
<tr>
<th>CHD* duration (years)</th>
<th>Carpal tunnel</th>
<th>Synovium</th>
<th>Flexor tendon</th>
<th>Bone cyst</th>
<th>Bursa</th>
<th>Synovial fluid</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>$\beta_2$ Microglobulin</td>
</tr>
<tr>
<td>12</td>
<td>+</td>
<td></td>
<td></td>
<td>Tibia</td>
<td>+</td>
<td>Strong</td>
</tr>
<tr>
<td>10</td>
<td>+</td>
<td></td>
<td>+</td>
<td>+</td>
<td></td>
<td>Weak</td>
</tr>
<tr>
<td>15</td>
<td>+</td>
<td>N†</td>
<td>+</td>
<td>N, carpi</td>
<td>+</td>
<td>Negative</td>
</tr>
<tr>
<td>14</td>
<td>+</td>
<td>N‡</td>
<td>N+</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>+</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td></td>
<td>§+</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*AM$_{\beta_{2m}}$=$\beta_2$ microglobulin related amyloid; CHD=chronic haemodialysis.

Only tissue sites in which deposits of AM$_{\beta_{2m}}$ were identified are indicated. $+=$ AM$_{\beta_{2m}}$ present; $N=$ necropsy findings; $t=$ shoulder, knee, hip, and lumbar apophysal joints; $t=$ shoulder and knee; $§=$ iliopsoas bursa excised seven years previously; this patient had no other features to suggest amyloidosis.
ropsy, deposits of AM\(_{p2m}\) were found in the synovium of shoulder, hip, knee, and carpal tunnel structures, including flexor tendon and retinaculum, but not median nerve. In the spine it was present in the synovium of lumbar apophyseal joints and in a degenerating lumbar disc but not in two adjacent, macroscopically normal, lumbar intervertebral discs. In general the amyloid was arranged in nodules (Fig. 3), which were not perivascular. Mononuclear cells, mainly lymphocytes, were numerous in the knee and shoulder but not the hip. There was also a moderate degree of synovial hyperplasia. Special stains for tissue deposits of iron showed moderate increases in synovial iron stores in the knees, a little in the shoulder, and none in other structures infiltrated by AM\(_{p2m}\), including hips, lumbar apophyseal joints, intervertebral disc or synovium of carpal tunnel, and flexor tendons. Synovial iron deposits were associated with a mononuclear infiltrate. No amyloid was identified in any other major organ.

A second patient, who had been dialysed for six years, had synovitis and restriction of shoulders and mild, clinically insignificant symptoms of CTS. At necropsy, six months after clinical review, AM\(_{p2m}\) was identified in the flexor retinaculum, median nerve, and synovium of shoulder and knee. No iron deposits or mononuclear cells were seen in any of these sites.

Necropsy was performed on three other subjects. Two had received haemodialysis for two years and one had had peritoneal dialysis for five months followed by two renal grafts lasting eight months before death. No amyloid was found in synovium from the carpal tunnel or adjacent structures, shoulders, hips, or knees. There had been no signs or symptoms of arthropathy or CTS during life.

Limited necropsy material was obtained from the finger of a sixth patient who had died two years after commencing CHD. The only articular abnormality in life was erosive nodal osteoarthritis affecting the distal interphalangeal joints. No amyloid deposits were found in the synovium or periarticular structures of an affected distal interphalangeal joint.

**Synovial fluid**

Synovial fluid obtained from the knees or shoulders of patients with chronic joint swelling was usually of high viscosity with small numbers of mononuclear cells. In three patients with histologically proved AM\(_{p2m}\) in synovial or carpal tunnel structures the synovial fluid sediment was positive on immunoperoxidase staining for \(\beta_2\) microglobulin (Table 6). In a further seven patients synovial fluid deposits also gave positive staining with immunoperoxidase. The staining pattern was usually fibrillar, but occasional amorphous clumps of AM\(_{p2m}\) were seen in 6/10 patients. Only one was strongly positive with

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![Fig. 3](http://ard.bmj.com/)

**Fig. 3** Nodular synovial deposit of \(\beta_2\) microglobulin related amyloid demonstrated by immunoperoxidase.
Congo red, however, three were weakly positive, and five were negative. Control studies using synovial fluid from patients with rheumatoid arthritis gave negative results using both techniques.

**BIOCHEMISTRY**

*Serum $\beta_2$ microglobulin*

Serum $\beta_2$ microglobulin concentrations before dialysis, which were available from 91/95 patients, were markedly raised (normal range 1–3 mg/l) and remained approximately constant with increasing duration of dialysis (Fig. 4). $\beta_2$ Microglobulin concentrations rose on average by about 4 mg/l during dialysis (data not shown), returning to predialysis concentrations between each session.

Within each cohort patients with chronic synovitis, CTS, or bone cysts did not have higher concentrations of serum $\beta_2$ microglobulin than those without these features (Table 7).

*Serum ferritin*

Serum ferritin concentrations were measured as an index of total iron stores and were available for 92/95 patients. Patients with large joint chronic synovitis had a wide scatter of serum ferritin concentrations, which clearly did not differ from those without synovitis (Fig. 5). Of the six patients with radiological evidence of large joint cartilage erosion without osteoarthritis, only one had modestly raised serum ferritin (340µg/l).

The relation between increased serum ferritin and
the development of osteoarthritis in large joints or small joints of the hands was also examined, but no correlation was found (Fig. 6). Many patients had substantially raised serum ferritin concentrations (>1000 μg/l) without any evidence of osteoarthritis, synovitis, or other arthropathy. Similarly, there was no correlation between raised ferritin concentrations and either CTS or bone cysts.

Discussion

Despite extensive reporting the epidemiology and aetiology of rheumatic disease in patients receiving CHD remain poorly defined. In this study of 95 patients receiving CHD the prevalence, clinical and radiological features of rheumatic disease, and its relation to AM2zm deposition, hyperparathyroidism, duration of dialysis, and various biochemical parameters were examined.

A striking finding was the high and increasing prevalence of symptoms and signs of joint disease with increasing duration of dialysis. The overall clinical picture was of a chronic arthropathy with synovitis or restriction, or both, affecting predominantly shoulders, knees, hips, and, to a lesser extent, elbows, wrists, and spine. Other important associations of this arthropathy included CTS, bone cysts, and, less frequently, flexor tenosynovitis of fingers. In a few patients haemarthrosis was seen. This clinical picture is consistent with previous reports, though hip involvement was more frequent than had been appreciated. Histological studies in life and at necropsy confirmed that deposition of AM2zm is a major cause of CTS and large joint swelling, and that AM2zm is a major constituent of bone cysts. It may also be responsible for the high prevalence of hip restriction. The deposition of AM2zm is clearly related to duration of dialysis and it was not found at necropsy in the synovium and carpal tunnel of three patients who had received dialysis for two years or less. Deposition of AM2zm may be accelerated at sites of inflammation as indicated by its presence in an iliopsoas bursa after three years of CHD. Seven years later this patient had no other evidence of amyloidosis.

Chronic swelling and restriction of large joints was not always accompanied by significant radiological evidence of joint damage. Of those with clinical abnormalities of knees, shoulders, and hips, 5/12, 5/16, and 10/17 patients respectively had no significant radiological abnormality. The main radiographic abnormalities identified included rotator cuff destruction, osteoarthritis with loss of cartilage, and, in a few, loss of articular cartilage without osteoarthritis.

The histological and synovial fluid examinations which were performed, taken in conjunction with other reported synovial histology, suggest that the synovial swelling in most patients with dialysis related arthropathy is due to massive infiltration by AM2zm. In the absence of synovial iron deposits no cellular infiltrate was seen. It cannot be assumed that synovial AM2zm is necessarily a cause of joint damage as about half of the patients with clinical or histological evidence of synovial AM2zm had no radiological abnormalities. Furthermore, the prevalence of joint lesions, such as cartilage erosion and severe rotator cuff disease, which have been attributed to AM2zm, were a function of age rather than duration of dialysis. Thus synovial deposition of AM2zm in damaged joints may simply be a secondary phenomenon, which is more likely to occur in diseased or inflamed tissue, rather than a primary aetiological event.

On the other hand, subchondral deposits of AM2zm may be directly responsible for joint damage. AM2zm deposits in subchondral bone have been reported to precede articular space narrowing by some years; destruction of articular cartilage was observed in one of three patients with a subchondral cyst in a large joint (Fig. 1c). Damage in this joint was localised to the immediate area over the deposit while the adjacent joint space appeared to be preserved. Degenerative changes were also noted in a metacarpophalangeal joint adjacent to a large phalangeal cyst. The mechanism of destruction is not certain but may simply be the mechanical effects of an enlarging deposit.

The possible role of hyperparathyroidism in causing large or medium joint damage was also examined, but no correlation was found between active hyperparathyroidism and either large or medium joint osteoarthritis, rotator cuff disease, or synovitis. Ligamentous laxity and collapse of subchondral bone, leading to destructive joint lesions and secondary osteoarthritis, can occur, however, in patients with primary or secondary hyperparathyroidism. Thus previous episodes of hyperparathyroidism may also be important in the aetiology of peripheral joint attrition.

Synovial iron deposition secondary to iron overload or recurrent haemarthrosis may also be an important factor in the pathogenesis of joint damage. Although there was no association between total iron stores, as measured by serum ferritin concentrations, and arthropathy, in one necropsy study a relation was noted between the amount of synovial iron, the number of infiltrating mononuclear cells, and the extent of joint damage. Caution is needed in interpreting these data as the iron and cellular infiltrate may simply be secondary to recurrent bleeding in a damaged joint.

Further prospective studies are needed to deter-
mine more precisely the relative contributions of synovial or subchondral deposits of AMβ2m, hyperparathyroidism, synovial iron deposits, and age to the pathogenesis of large and medium joint arthritis in patients receiving CHD.

Symptoms of pain and stiffness in the hands were frequent and increased markedly with increasing duration of dialysis. In four patients this was attributed clinically to nodular tenosynovitis, which in two was due to AMβ2m deposition within the flexor tendons. Osteoarthritis was also responsible for symptoms in some patients, but in this series erosive osteoarthritis was correlated with age rather than duration of dialysis. In two patients it occurred at an unusually young age, suggesting that factors associated with renal failure may contribute. No AMβ2m was identified in an affected distal interphalangeal joint of a patient with erosive osteoarthritis at necropsy, confirming that amyloid plays no part in this condition. Most of the 14 patients with erosive azotaemic osteoarthropathy had no small joint symptoms, and the condition did not correlate with evidence of active hyperparathyroidism. This is concordant with a study by Sundaram et al but not with that by Rubin et al. Overall, the increasing frequency of small joint pain and stiffness with duration of dialysis was not satisfactorily accounted for by clinical signs of synovitis or deformity, or by radiographic evidence of joint space narrowing, osteoarthritis, or erosive azotaemic osteoarthropathy.

Pain, stiffness, and restriction of the spine were relatively common, and in a few symptoms were attributable to mild degenerative disc or facet joint disease. Two patients had destructive disc lesions with loss of adjacent end plate. The appearances were similar to the destructive spondyloarthropathy reported recently, which has been attributed in turn to eitherapatite or AMβ2m deposition. One of these two patients had extensive AMβ2m deposition in synovium, including apophyseal joints, and bone; AMβ2m also was found in the degenerate lumbar vertebral disc but not in adjacent macroscopically normal lumbar discs. This patient had previously had a parathyroidectomy and had a rugger jersey spine. Similar destructive spondyloptic lesions have been described in primary and secondary hyperparathyroidism and appear to be caused by collapse of juxta-articular bone. This may have been the mechanism in our two patients and in the others in whom this has been reported. As premorbid deposits of AMβ2m were absent from other macroscopically normal discs in the patient described above AMβ2m deposition is probably a consequence rather than a cause of disc degeneration.

AMβ2m was identified by immunoperoxidase in synovial fluids from several patients with suspected or histologically proved synovial AMβ2m. Control studies on synovial fluid from patients with active rheumatoid arthritis were negative. Synovial fluid from a patient with acute exacerbation of ankylosing spondylitis, who had recently started CHD treatment and is therefore not included in the survey, also gave negative results for AMβ2m. Attempts to verify that positive immunochromal results correlated with the presence of Congophilic material were largely unsuccessful. The failure to identify AMβ2m using Congo red may be due to the lack of sensitivity of this technique compared with immunochromal methods. Recently we treated a synovial fluid sample from a newly diagnosed patient with hyaluronidase in an attempt to ensure more complete recovery of fibrils during centrifugation. This specimen gave positive results with both immunoperoxidase and Congo red techniques. It seems likely that failure to obtain Congo red staining in sediments which are immunoreactive for β2 microglobulin is due to false positive results with the immunochromal technique; for example, soluble β2 microglobulin may simply have precipitated as an amorphous deposit. This technique requires further evaluation before it can be accepted as diagnostically useful.

We also examined the relation between serum β2 microglobulin and the AMβ2m related syndrome. Contrary to some reports, but in agreement with others, we found little or no rise in serum β2 microglobulin concentrations with increasing duration of dialysis. Also as reported previously, we found a transient rise in serum β2 microglobulin during each dialysis and no correlation between the development of the AMβ2m related syndrome and serum β2 microglobulin concentrations. The final total body pool of β2 microglobulin—that is, soluble and insoluble—is presumably a more important factor than serum β2 microglobulin in determining the development of the AMβ2m related syndrome.

An important question is why some patients do not develop dialysis related arthropathy and its associated features even after many years of CHD. Five of the 14 patients in cohort C had no features—that is, CTS, bone cysts, or synovitis, to suggest that they had the AMβ2m related syndrome. The mean (SD) age of these five patients (38.6 (4.76) years) was significantly less than that of the other nine (59.7 (10.71) years) (p<0.01; Student’s unpaired t test). This trend was also apparent in cohorts A and B but did not reach statistical significance. Thus age may be an important factor determining predisposition to the development of amyloidosis.
In summary, at least three clinically distinct subgroups of patients with arthropathy were identified—patients with the AMβ2m related syndrome of synovial swelling, tenosynovitis,CTS, and bone cysts; a second group with erosive azotaemic osteoarthropathy; and a third group with degenerative joint disease affecting small, large, or axial joints.

A number of important conclusions may be drawn concerning the aetiology of these conditions. Firstly, AMβ2m deposition increases with duration of dialysis and causes synovial swelling, nodular tenosynovitis, bone cysts, and CTS; in patients receiving long term (>10 years) CHD, however, the risk of developing AMβ2m was related to age. Deposition of AMβ2m in subchondral bone, but not synovium, causes joint damage, and AMβ2m may be more prone to deposition in synovium already damaged by other pathological processes. In the absence of synovial iron deposition synovial AMβ2m is not associated with an inflammatory infiltrate. Although the data are inconclusive, hyperparathyroidism may have a role in the pathogenesis of destructive spondyloarthropathy and perhaps of the degenerative disease of peripheral joints in some patients. The failure to confirm a correlation between hyperparathyroidism and erosive azotaemic osteoarthropathy may be due to difficulties in the accurate ascertainment of hyperparathyroidism in patients receiving CHD. No correlation was found between serum ferritin and arthritis, and the significance of synovial iron deposition is unclear.

As some of the features of dialysis related arthropathy may be ameliorated by alterations in dialysis technique or other therapeutic intervention further well designed prospective studies are urgently needed to clarify the pathogenetic mechanisms responsible for joint damage in these patients.

We thank Professor Barry Vernon-Roberts for reporting results of bone histology, Mr David Gove for performing electron microscopy studies, and Dr S C Milazzo for helpful criticism of this manuscript.

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Ann Rheum Dis 1989 48: 409-420
doi: 10.1136/ard.48.5.409

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