Amyloid arthropathy in patients undergoing periodical haemodialysis for chronic renal failure: a new complication

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Summary Seven patients (five male and two female) with chronic renal failure (CRF) treated by periodical haemodialysis presented with swelling and effusion of more than three months' duration in knees (four bilateral), shoulders (two, one of them bilateral), elbow (one), and ankle (one). Four had a carpal tunnel syndrome both clinically and electromyographically (three bilateral). All patients had hyperparathyroidism secondary to their CRF, which was not due to amyloidosis in any of them. The dialysis duration period varied from five to 14 years, with an average of 8·6 years. Amyloid deposits (Congo red positive areas with green birefringence under polarizing microscopy) were shown in six of the seven synovial biopsy specimens of the knee, in five of the sediments of the synovial fluids, and in specimens removed during carpal tunnel syndrome surgery. No amyloid was found in the biopsy specimen of abdominal fat of six of the patients. The finding of amyloid only in the synovial membrane and fluid, and carpal tunnel, its absence in abdominal fat, and the lack of other manifestations of generalised amyloidosis (cardiomyopathy, malabsorption syndrome, macroglossia, etc.) and of Bence Jones myeloma (protein immunoelectrophoresis normal) raises the possibility that this is a form of amyloidosis which is peculiar to CRF treated by periodical haemodialysis.

Key words: amyloidosis, synovial amyloidosis, synovial fluid, carpal tunnel syndrome.

Various musculoskeletal disorders have been described in patients with chronic renal failure (CRF) treated by periodical haemodialysis. Among these are synovitis and bursitis due to sodium urate, calcium pyrophosphate, hydroxyapatite, and calcium oxalate microcrystals, bone and joint infections, osteonecrosis, carpal tunnel syndrome, soft tissue and vascular calcifications, and metabolic bone disease (hyperparathyroidism, osteomalacia, osteoporosis, osteosclerosis).

Other alterations have been described: microtraumatic olecranon bursitis caused by the arm position during haemodialysis, unexplained 'dialysis cysts' (small cystic lesions in the hand and wrist bones), digital clubbing in one or more fingers induced by anoxia distal to the fistula, and aneurysm at the site of the shunt, with or without calcification.

Our relationship with the Nephrology Department in the follow up of CRF patients in a periodical haemodialysis programme enabled us to identify several patients with swelling and effusions that could not be explained by the above-mentioned causes.

The clinical picture in the first patient studied (case No 1) was a chronic swelling with large effusion in the knees and shoulders (similar to the 'shoulder pad sign'), which suggested the possibility of amyloid arthropathy.

Materials and methods

CRF patients undergoing periodical haemodialysis, with persistent swelling and effusion in the knee...
Some of the samples were treated with potassium permanganate by Wright's technique, and in six out of seven patients a needle aspiration of abdominal fat was performed.

Results

The group studied included five men and two women aged from 33 to 62 years (mean age 50) (Table 1). The aetiology for their chronic renal failure was: glomerulonephritis (two), backflow hydronephrosis (one), nephroangiosclerosis (one), and unknown (three). No patient had renal amyloidosis.

The dialysis duration period varied from five to 14 years, with an average duration of 8-6 years.

Clinical examination of the patients showed swelling and effusion in the major joints: knees (seven patients, four of them bilateral), shoulders (two patients, one of them bilateral), ankle (one patient), and elbow (one patient). The swellings were accompanied by minor pain, their onset being therefore difficult to ascertain; during our observations their duration varied between six months and two years. Two patients presented with an acute crisis in knees, shoulder, and elbow, with very 'inflammatory' fluids during the crisis, which caused the patients to be admitted and a culture of the fluid to be undertaken

Table 1 Clinical features of the patients studied

<table>
<thead>
<tr>
<th>Patient No</th>
<th>Sex</th>
<th>Age</th>
<th>Approx. CRF age</th>
<th>Dialysis (years)</th>
<th>Joint signs and duration</th>
<th>Carpal tunnel syndrome</th>
<th>Radiology</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>M</td>
<td>59</td>
<td>49</td>
<td>9</td>
<td>Persistent swelling knees/shoulders more than one year. Acute crises shoulders, knee, elbow</td>
<td>Yes. Bilateral. Operated both sides</td>
<td>Geodes in shoulders. Hyperparathyroidism</td>
</tr>
<tr>
<td>2</td>
<td>M</td>
<td>48</td>
<td>?</td>
<td>10</td>
<td>Persistent swelling knees and ankle, two years</td>
<td>No</td>
<td>Geodes in shoulders, knees, and pubis. Hyperparathyroidism</td>
</tr>
<tr>
<td>3</td>
<td>M</td>
<td>47</td>
<td>37</td>
<td>5</td>
<td>Persistent swelling knees and right shoulder, six months</td>
<td>Yes. Bilateral. Operated both sides</td>
<td>Microgeodes in hands. Hyperparathyroidism</td>
</tr>
<tr>
<td>4</td>
<td>M</td>
<td>54</td>
<td>?</td>
<td>8</td>
<td>Persistent swelling left knee, one year. Right shoulder joint pain</td>
<td>Yes. Bilateral. Operated both sides</td>
<td>Geodes in knees, right femur, and right shoulder. Hyperparathyroidism</td>
</tr>
<tr>
<td>5</td>
<td>F</td>
<td>62</td>
<td>50</td>
<td>9</td>
<td>Persistent swelling both knees, one year. Shoulder joint pains</td>
<td>Yes. Right operated</td>
<td>Patella subluxation bilateral. Vertebral osteodystrophy</td>
</tr>
<tr>
<td>6</td>
<td>M</td>
<td>50</td>
<td>?</td>
<td>5</td>
<td>Persistent swelling both knees, one year. Acute crises both knees</td>
<td>No</td>
<td>Arthrosis right knee (meniscectomy)</td>
</tr>
<tr>
<td>7</td>
<td>F</td>
<td>33</td>
<td>19</td>
<td>14</td>
<td>Persistent swelling left knee, six months. Joint pains in shoulders, ankles, right knee</td>
<td>No</td>
<td>Hyperparathyroidism</td>
</tr>
</tbody>
</table>
to check the possibility of an infection in the joints, but the results were negative. Between the crises there was persistence of the effusions with minor pain and low cellularity. Four patients complained of pains in shoulders, with no visible swelling at any time.

Four patients presented with a carpal tunnel syndrome both clinically and electromyographically, bilateral in three of them.

Radiological signs of hyperparathyroidism or osteodystrophy, or both were seen in all the cases and were intense in three of them.

Three patients showed moderate or severe cystic lesions in shoulders (three), knees (two), and the superior branch of the pubic symphysis (one). One patient showed microgeodes, and another patient developed a bilateral external subluxation of the patella.

The blood biochemistry profile showed CRF and the classical alterations of secondary hyperparathyroidism in all patients. Serum protein electrophoresis was normal; serum immunoglobulins levels (IgA, IgG, IgM) were normal in all patients except for one with an IgG of 1760 mg/100 ml (17.6 g/l) (normal

<table>
<thead>
<tr>
<th>Patient No</th>
<th>Synovial fluid</th>
<th>Amyloid deposits median nerve</th>
<th>Fat</th>
<th>Synovial membrane</th>
<th>Synovial fluid</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Non-inflammatory, Inflammatory in crises</td>
<td>Positive. Annular ligament (1) synovium (1)</td>
<td>Negative</td>
<td>Positive. Mesothelial hyperplasia</td>
<td>Positive. Shoulder(s) knee(s)</td>
</tr>
<tr>
<td>3</td>
<td>Non-inflammatory</td>
<td>Positive. Perineural region (2)</td>
<td>Negative</td>
<td>Positive. Fibrous connective tissue</td>
<td>Positive. Knee(s)</td>
</tr>
<tr>
<td>5</td>
<td>Non-inflammatory</td>
<td>Positive. Penneural region (2)</td>
<td>Negative</td>
<td>Positive. Non-specific chronic synovitis</td>
<td>Positive. Knee(s)</td>
</tr>
<tr>
<td>6</td>
<td>Slight inflammatory, Severe in crises</td>
<td>—</td>
<td>Negative</td>
<td>Negative. Mesothelial hyperplasia</td>
<td>Not done</td>
</tr>
</tbody>
</table>

Fig. 1 Synovial tissue: (a) amorphous material deposition Congo red positive; (b) showing intense green birefringence under polarising microscopy. (Congo red, ×190.)
values 600–1500 mg/100 ml (6–15 g/l)). Immunoelctrophoresis showed no qualitative abnormality of immunoglobulins nor the presence of light chains. These results excluded the presence of a Bence Jones myeloma in our patients.

The synovial fluid was 'non-inflammatory' in six patients (<2000 leucocytes/mm\(^3\) (2×10\(^9\)/l)), and the remaining fluid showed moderate inflammatory characteristics (around 5000 cells/mm\(^3\) (5×10\(^9\)/l)). Two patients who suffered an occasional acute crisis showed very cellular fluids during these crises, fluctuating between 8000 and 103 000 cells/mm\(^3\) (8–103×10\(^9\)/l) with an average of 30 000 leucocytes/mm\(^3\) (30×10\(^9\)/l).

No crystals were observed in any of the fluids, either in the unstained examination or after alizarin red staining.

The histological study (Table 2) of the synovial biopsy specimens showed amyloid deposits (Figs 1a and 1b) in six patients (Congo red positive areas with green birefringence); the other findings varied greatly: slight mesothelial hyperplasia (four patients, one of them amyloid negative), non-specific chronic synovitis (one), fibrous connective tissue (one), and normal (one).

The presence of amyloid in the synovial fluid sediment (Fig. 2) was proved in five of the six patients studied (fluid from the knees in six occasions, from the shoulder in one); the negative fluid showed amyloid in the synovium. No fluid was studied in the seventh patient (No 6) with an amyloid negative synovial biopsy specimen.

This patient had a clinical picture similar to that of the others, with swelling and effusion in both knees for a little over one year, treated with aspiration and injections of a depot corticosteroid every three to four months. The patient had been referred from another hospital, and subsequent checks for the presence of amyloid in the effusion were not possible.

Surgery was performed on four of the patients with a carpal tunnel syndrome (two of them bilateral), showing amyloid deposits in all of them (in the annular ligament in two, in the perineural region in two, and in the synovium in another).

A biopsy specimen of the abdominal fat was taken from six patients and proved amyloid negative in all of them.

Potassium permanganate staining was positive in all articular fluid sediments and in the perineural region for the only patient who had an operation for carpal tunnel syndrome in which it was studied.

**Discussion**

Six of the patients had synovial amyloidosis, four of them with a carpal tunnel syndrome, a frequent location of the so-called primary amyloidosis, but without any other manifestations of generalised amyloidosis (cardiomyopathy, malabsorption, macroglossia, etc.).

In patient No 6 the synovial biopsy specimen probably did not include any affected synovium, or the aetiology for this case might have been different. The positive results from the treatment of the samples by Wright’s technique pointed towards this classification, but the lack of amyloid in the biopsy specimen of the abdominal fat, negative in the six patients in which it was sought and which is usually positive in primary and generalised amyloidosis, did not allow us to classify these patients as having this type of amyloidosis.

The protein immunoelctrophoresis normal values without light chains confirmed that our patients did not suffer from a Bence Jones myeloma.

All the patients had hyperparathyroidism secondary to their chronic renal failure, and this was not due to amyloidosis in any of them.

The literature to date has described patients with idiopathic generalised amyloidosis associated with a Bence Jones myeloma. In our patients the clinical picture was one of chronic synovitis affecting mainly knees and shoulders, showing amyloid deposits in

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**Fig. 2** Synovial fluid centrifugate. Multiple deposits of Congo red positive material with green birefringence. (Polarising microscopy, Congo red, ×100.)
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synovial membrane and in the carpal tunnel, but without deposition in the abdominal fat or other manifestations of generalised amyloidosis; for these reasons we think that we are probably dealing with a form of amyloidosis which is peculiar to CRF treated with periodical haemodialysis.

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References


Book review


Laboratory analysis of synovial fluid provides a firm diagnosis in only two types of arthritis: microbial infections and crystal synovitis. It can also identify a haemarthrosis and give some idea of the acuteness of synovial inflammation; but here the laboratory does not add very much to what is apparent from naked eye inspection of the fluid. The minimum requirements for the clinician dealing with rheumatic diseases are, therefore, facilities for the reliable identification of synovial crystals and infections.

Identification of synovial infections has so much in common with the requirements of other specialties that routine microbiology laboratories generally handle this adequately. Crystal identification, however, requires experience and hardware that is not so often available in routine laboratories. For this reason, many rheumatologists have found it necessary to equip themselves to carry out this investigation, and a generation of trainees is coming to regard this as an essential part of their experience.

An excellent account of all aspects of the examination of synovial fluids is included in the recent 10th edition of ‘Hollander’, this section being written by the editor himself—Dr Dan McCarty. Do we, therefore, need this new book on the same subject? I believe we do. This handy, slim volume of about 100 pages is well set out and includes excellent illustrations. Its place is beside the polarising light microscope in the outpatient department, and on the shelf of the clinical pathology laboratory. It needs to be within easy reach of both the beginner who cannot remember the optical properties of common crystals and also the more experienced, who will find useful information about such matters as the identification of injected corticosteroid material and other uncommon contaminants. There is a fund of other useful information about all aspects of synovial analysis.

I was surprised at two pieces of advice given. The first is to use an anticoagulant when taking fluid for crystal examination; I find that flecks of clot are useful to focus on and, by trapping crystals, provide an enriched area in which to search. Secondly, the author recommends using a high power oil immersion lens when studying ‘wet’ preparations for crystals. This may increase the sensitivity of the test, but it must also make it more complicated and time-consuming.

However, in general the advice given is clear, comprehensive, and essentially practical (the author’s advice on arthrocentesis includes keeping a small artery forceps handy for removing needles when switching syringes or extracting the distal fragment of a broken needle!). I recommend it.

HARRY CURREY
Editor


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