Mixed systemic amyloidosis in a patient receiving long term haemodialysis

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Abstract
A 64 year old woman had been receiving haemodialysis for 11 years. She had a history of chronic renal failure, caused by probable chronic pyelonephritis, and dialysis arthropathy. She died of acute pulmonary oedema and haemorrhage. At necropsy, histological, immunohistochemical, and ultrastructural studies showed widespread visceral deposits of \( \beta_2 \)-microglobulin (\( \beta_2 \)-M) and AA amyloid.

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Beta\(_2\)-microglobulin (\( \beta_2 \)-M) amyloidosis that predominantly affects perineural and periarticular structures, joints, and bone is common in patients receiving long term haemodialysis.\(^1\) Systemic deposits of \( \beta_2 \)-M amyloid have also been reported. Initially, these findings were considered clinically unimportant.\(^2\) Subsequently, however, some investigators showed that extra-articular deposits of \( \beta_2 \)-M amyloid can cause serious clinical problems.\(^3,4\) On the other hand, non-\( \beta_2 \)-M type amyloidosis also appears in patients receiving haemodialysis, and this is generally attributed to concomitant disease.\(^5\)

We report a patient who had been receiving haemodialysis for 11 years with dialysis associated amyloid arthropathy in whom a post mortem examination showed systemic deposits of AA and \( \beta_2 \)-M amyloid.

Case report
A 64 year old woman with no history of tuberculosis, multiple myeloma, malignancy, osteomyelitis, bronchiectasis, chronic inflammatory bowel disease, or articular inflammatory disease, was diagnosed with end stage renal failure for probable chronic pyelonephritis in 1978. She required haemodialysis from December 1979. In March 1988 she received right-sided carpal tunnel decompression for carpal tunnel syndrome. A biopsy specimen obtained at the operation showed amyloid deposits of \( \beta_2 \)-M type. During the same year a parathyroidectomy was performed because of secondary hyperparathyroidism. Two abdominal fat biopsy specimens taken in 1988 and 1989 showed no amyloid deposits. In 1990 she had recurring infections of the arteriovenous fistule for seven months. She died that year because of acute pulmonary oedema and intestinal bleeding.

The principal macroscopic findings were: mild cardiomegaly with left myocardial hypertrophy; calcification of the mitral valve and aortic cusps; adhesive fibrous pericarditis; scleratrophic kidneys with cystic dialysis transformation; haemorrhagic colitis; pulmonary oedema; and inactive bronchiectasis in the medium lobe of the right lung. No malignancies or tuberculosis were found.

Microscopic examination of tissues fixed in 10% buffered formalin, disclosed abundant amorphous eosinophilic material which stained with alkaline Congo red and showed apple-green dichroism under polarised light. The amyloid had infiltrated the vessel walls of visceral organs (gastrointestinal tract, liver, spleen, pancreas, heart, urinary bladder, and kidneys) and skin; muscular layers of the gastrointestinal tract and urinary bladder; and myocardium, pericardium, and peritoneum. Amyloid deposits were particularly heavy in the auricular myocardium.

In the vessel walls amyloid was found in the muscular layer and in the subendothelial space where it appeared as nodules protruding into the vascular lumen. An intense foreign body giant cell reaction was found around the scattered submesothelial nodules.

Immunohistochemical studies using the avidin-biotin complex method were performed on serial sections of the paraffin wax embedded samples of stomach, ventricular and auricular myocardium, pericardial and peritoneal nodules. Monoclonal antibodies to amyloid \( A (AA) \), and polyclonal \( \beta_2 \)-M, pre-albumin, \( \kappa \) and \( \lambda \) light chains (Dako) were applied. For positive controls, substitution of the antibodies with TRIS-buffered saline, and alkaline Congo red staining of tissue samples were also used.

Only anti-AA and \( \beta_2 \)-M antibodies reacted positively with the amyloid. In the stomach both antibodies stained the amyloid of the muscularis propria with anti-AA showing a more diffuse pattern of staining than anti-\( \beta_2 \)-M; in suberosal nodules \( \beta_2 \)-M was mainly observed in the periphery; AA appeared in the core (figure). In the vessels of all samples anti-\( \beta_2 \)-M seemed to be generally confined to the subendothelial amyloid, and anti-AA stained the other layers of the wall. In the myocardium the amyloid reacted positively only with anti-AA, while in the auricular myocardium both antibodies appeared intermingled.

Electron microscopic studies carried out on formalin fixed auricular myocardium showed interstitial curvilinear microfibrils...
arranged in bundles characteristic of β2-M amyloid, in addition to crossed straight microfibrils arranged haphazardly, characteristic of the other types of amyloid.  

Discussion

Dialysis amyloidosis seems to have a predilection for the osteoarticular system, but in some instances visceral amyloid deposits can cause serious clinical complications. In our patient clinical manifestations of visceral amyloidosis were absent, at least until the episode of heart failure that led her to death.

It is worth pointing out several findings in this case. Firstly, the results of the histological, immunohistochemical, and ultrastructural studies showed the presence of systemic deposits of both amyloids (β2-M and AA) which happens only rarely. Secondly, both amyloids showed a distinctive tissular distribution, with β2-M being preferentially located in the subendothelium of the vessels, as noted before, and surrounding the AA amyloid deposits in the mesothelial nodules. Finally, an intense giant cell reaction was present around the β2-M amyloid deposits. Giant cell reaction to β2-M, a common finding in the synovium, has occasionally been described in visceral deposits.

Several factors, such as chronic pyelonephritis, bronchiectasis, repetitive infections of the arteriovenous fistule, and haemodialysis could explain the presence of AA amyloid in this case, though it is difficult to ascribe its development to one of them. It seems unlikely that bronchiectasis could account for AA amyloidosis because of the absence of previous history and its inactive character. In a review of 45 consecutive necropsies with bronchiectasis we found AA amyloidosis in four cases, all of them with clinical and pathological evidence of active disease (unpublished data).

As regards chronic pyelonephritis, AA amyloidosis has scarcely been reported in the absence of stag horn calculi or spinal cord injury. No conditions were present in this case.

The patient had had repeated infections of the arteriovenous fistule for seven months before her death, but it is difficult to consider those infectious episodes as the cause of AA amyloidosis in such a short time.

A constant synthesis of AA protein induced by the leucocyte pyrogen produced by haemodialysis might be another possible mechanism for the development of AA amyloidosis in this patient.

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