Mobile intracardiac calcinosis: a new risk of thromboembolism in patients with haemodialysed end stage renal disease

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Abstract
Cardiac calcinosis is a common complication of end stage renal disease. A newly observed risk of thromboembolism is reported in four patients with mobile cardiac calcinosis, treated with long term dialysis. Rapidly growing mobile calcification was confirmed by echocardiography. Each patient had an imbalance in serum calcium × inorganic phosphate (Ca × P product > 50); this imbalance could not be treated due to the sudden death of the patient or the need for surgical resection to prevent recurrent cerebral thromboembolism. Histological examination revealed intracardiac calcinosis in three cases, and each case showed haemodialysis hypoparathyroidism (intact PTH < 160 pg/ml). Thromboembolism in such cases is rare, however it indicates a need for cautious echocardiographic monitoring in end stage renal disease patients with an uncontrolled Ca × P product. (Heart 1999;82:638–640)

Keywords: cardiac mass; intracardiac mobile calcinosis; haemodialysis hypoparathyroidism; thromboembolism; end stage renal disease

Abnormal calcification of the cardiovascular system—for example, mitral annular calcinosis, calcified aortic root, and arterial calcification, is a well known complication of end stage renal disease treated with long term haemodialysis, and such complications often result in fatal cardiac complications such as valvar heart disease, congestive heart failure, and obstructive and atherosclerotic arterial disease. The serum calcium × inorganic phosphate product (Ca × P product) is thought to have a partial effect on systemic, but not cardiac, calcinosis in patients treated with haemodialysis using high performance membrane which controls for serum calcium inorganic phosphate levels. Furthermore, little consideration has been given to the possibility of thromboembolism in intracardiac calcified deposits in end stage renal disease.

This paper discusses four cases of rare mobile pseudocardiac tumours, which were histologically proved to be intracardiac calcinosis, and is the first report of intracardiac mobile calcinosis with thromboembolism in haemodialysed end stage renal disease patients.

Case 1
A 70 year old woman with interstitial nephritis due to recurrent pyelonephritis had been treated with dialysis since 1991. Since 1995, she had effort angina and was admitted in early 1997 for examination of this condition. Stress myocardial scintigraphy revealed reversible ischaemia in the anteroseptal and inferior region, and antianginal drugs, including β blockade, enabled her to reach an exercise capacity of 5.0 METs. Haemodialysis was performed with bicarbonate dialysate three times a week.

In February 1998, the patient was again admitted for echocardiography, which showed a crotchet shaped mobile interventricular mass (fig 1). Laboratory examination revealed renal anaemia with haemoglobin 73 g/l. Serum calcium was 10.5 mg/dl, phosphorus was 6.0 mmol/l, and aluminium was 593 nmol/l. After treatment with a supplement of CaCO3 (2.6 mmol/l (10.5 mg/dl) × P (2.3 nmol/l (7.0 mg/dl))) was 73.6. Serum parathyroid hormone (intact PTH) was 16.9 pg/ml, magnesium was 0.7 mmol/l, vitamin D was 26 nmol/l, and aluminium was 593 nmol/l. After treatment with a supplement of CaCO3 (3.0 g daily) and active vitamin D3 (alfacalidol 0.25 µg daily), Ca × P product dropped to 48.4. However, renal anaemia worsened, and the patient developed angina at night with dyspnoea. In June 1998, she felt orthopnoea at night, which could not be relieved with intravenous glycerol trinitrate. She died four hours later from cardiogenic shock.

Necropsy revealed a white steak-like mass attached to the membranous portion of the interventricular septum. Microscopic evaluation showed typical tumoral calcinosis and continuous calcification of the mitral annulus and the aortic root, which was partially connected to interventricular septum where the tumour was attached. Fresh massive myocardial infarction was also confirmed, with triple vessel disease.

Case 2
A 66 year old woman with renal failure due to renal tuberculosis after nephrectomy had been

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Intracardiac mobile calcinosis in end stage renal disease

We report four cases of mobile cardiac calcinosis accompanied by mitral annular and subaortic calcification; this is also the first report of cardiac tumoral calcinosis as a cause of thromboembolism in end stage renal disease. These cases illustrate a very rare complication, which is important in patient prognosis in end stage renal disease.

It is not usual to perform frequent echocardiographic evaluation in end stage renal disease. We cannot comment on the precise incidence of intracardial tumoral calcinosis in our study, but we speculate that the incidence of mobile calcinosis is not high, as the echocardiographic database on end stage renal disease in our three hospitals has only 300 cases. Detection of mitral annular calcinosis and calcification of the aortic root by echocardiography is not difficult, but annular calcinosis can mimic an intracardiac mass. In our cases, early detection of mobile calcinosis had failed even with an interval of 6–12 months between evaluations. A further difficulty with diagnosis is how to differentiate mobile calcinosis from primary cardiac tumours or intracardiac thrombi; dense continuity from the mitral or calcified aortic root and homogeneous high echogenic characteristics can be helpful in such cases.

Case 1
A 55 year old woman with diabetic nephropathy had been treated with continuous ambulatory peritoneal dialysis for four years. Echocardiography (fig 1) in late 1996 revealed a mobile left atrial mass attached to the base of the posterior left atrial wall, and she was admitted for further evaluation. Due to the rapid growth of the mass, we recommended surgical resection, but the patient rejected this suggestion. The patient also had renal anaemia with haemoglobin 53 g/l, and the Ca (2.7 mmol/l (11.0 mg/dl)) × P (1.6 mmol/l (5.1 mg/dl)) product was 56.1; her intact PTH level was 17.1 µg/ml. Four months later she died suddenly. Necropsy revealed destruction of the intra-atrial mass, which had been attached to the posterior left atrial wall, and embolism in the cerebrovascular system.

Case 2
A 63 year old man with chronic glomerulonephritis had been haemodialysed for eight years. One year before admission, he had felt dyspnoea on effort. When atrial fibrillation became his dominant rhythm, his exercise was limited to New York Heart Association (NYHA) functional class III. He had been taking calcium bicarbonate 3.0 g and active vitamin D, 0.50 µg daily for eight years.

The patient was admitted in June 1998. Echocardiography revealed noticeable mitral annular calcinosis with left atria mobile tumours attached to the base of the anterior mitral leaflet; 1.1 cm² of functional mitral stenosis was also detected in the Gorlin mitral valve area. The Ca (2.9 mmol/l) × P (1.7 mmol/l) product was 62.0; the patient’s intact PTH was severely depressed at 9.0 µg/ml. Magnesium, vitamin D, and aluminum levels were all within normal range.

Figure 1  Case 1: transoesophageal echocardiography (TEE) showing a mobile intracardiac tumour (6 × 14 mm) with homogenous hyperechoic characteristics, attached to the membranous portion of interventricular septum with tumour. Case 2: echocardiography after cerebral thromboembolism, showing a heterogeneous high echoic mobile intracardiac tumour at the base of the posterior mitral leaflet (PML), and the rapid growth of the tumour (14.7 × 13.7 mm). Case 3: echocardiography showing rapidly growing intracardiac mass (14 × 12 mm) on posterior wall of left atrium (LA; mid-panel). Case 4: transoesophageal echocardiography showing a homogeneous high echoic mobile intra-atrial tumour located in anterior mitral valve (AMV) (8 × 12 mm). AML, anterior mitral leaflet; Ao, aorta; LV, left ventricle; MAC, mitral annular calcinosis; RV, right ventricle.
Systemic massive calcinosis on extraskeletal tissue is not a rare complication of end stage renal disease, especially in patients treated with long term haemodialysis, cardiac calcinosis, like mitral annular calcinosis and subaortic calcification, is also common. However, it is not clearly understood why the cardiac base is vulnerable to calcium deposits. In extraskeletal calcinosis, Ca−P deposits on the multifollicular fibrous wall are also seen frequently in the hyperkinetic joint capsule. The similarity of the histological and pathophysiological background of the target tissue may explain the deposit of fibrous tissue in the cardiac base. But the clinical significance of cardiac calcinosis is very different from joint calcinosis because the rapid progression of cardiac calcinosis is due to exposure to the bloodstream, which could produce a secondary thrombus formation. Furthermore, tissue resistance, which might limit the growth of the tumour, is very low and speed of growth in cardiac tissue could be too high. Finally, the early detection of cardiac calcinosis is more difficult than the detection of joint calcinosis.

Secondary hyperparathyroidism plays a key role in the cause of cardiac calcification, and a serum marker may be an increased level of inorganic phosphate and serum calcium, where the Ca × P product is < 50. Serum magnesium, vitamin D, and serum aluminum are all influential factors. In our cases, only the Ca × P product was high. Modification of treatment through the use of low calcium dialysate, high performance membrane, and a drug such as vinpocetine on the Ca × P product, gave some beneficial effects for systemic tumoral calcinosis, but not for cardiac calcinosis.

A further symptom of rapid cardiac calcinosis is hypoparathyroidism. Recently, Akizawa et al reported that patients on haemodialysis with hypoparathyroidism had a higher incidence of metastatic calcification, which resulted from the decreased reservoir function of bone for absorbed calcium and phosphate from the intestine. Moreover, generally administered calcium and active vitamin D could exacerbate hypercalcaemia, and might increase the risk of tumoral calcinosis in the cardiovascular system of patients with end stage renal disease and hypoparathyroidism. Patients with hypoparathyroidism are defined as having intact PTH < 160 pg/ml. Thus, all of our cases may have had dialysis related hypoparathyroidism, which could have contributed to the development of cardiac calcinosis, rather than the calcinosis being a result of a Ca × P product of < 50.

Nevertheless, it seems that anticoagulation treatment is essential for the prevention of secondary formed thrombi. Additionally, surgical resection should be considered to be a primary treatment, as mobile tumoral calcinosis in the cardiovascular system carries a high risk of thromboembolism of cardiac origin.

In conclusion, cardiac mobile tumoral calcinosis, a rare complication of end stage renal disease, carries a high risk of thromboembolism, especially when the patient has poorly controlled serum Ca × P product and haemodialysed hypoparathyroidism. A cautious echocardiographic evaluation could also be useful in the detection of mobile cardiac tumoral calcinosis.