Left ventricular morphology and diastolic function in uraemia: echocardiographic evidence of a specific cardiomyopathy

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Abstract

Objective—To see whether cardiac morphological and functional abnormalities in uraemic patients are determined by high blood pressure or if they are an expression of a specific cardiomyopathy.

Setting—City general hospital in Italy.

Subjects—35 uraemic patients receiving haemodialysis (17 men, 18 women; mean age 60±3 (11·2); mean duration of dialysis 52 months) were selected from the 64 patients in Venice who were receiving dialysis; subjects with diabetes, haemochromatosis, valvar dysfunction, regional dyskinesias, and pericarditis were excluded. 19 control normotensive subjects (6 men and 13 women), matched for age.

Main outcome measures—Echocardiographic measurements of left atrium, left ventricular end diastolic and end systolic volume, aortic root diameter, posterior wall and interventricular septum thickness, left ventricle mass index, and ejection fraction in controls and in patients according to whether they were normotensive (five men, eight women) or hypertensive (12 men, 10 women) on 48 hour ambulatory monitoring; left ventricular diastolic function by Doppler ultrasonography.

Results—Mean systolic and diastolic pressures, daytime systolic and diastolic pressures, and night time systolic and diastolic pressures were significantly higher in the hypertensive patients than in the normotensive patients. The normotensive patients had similar blood pressures to the controls. Left ventricular mass correlated significantly with the mean diastolic pressure and mean night time systolic and diastolic pressures. Parathyroid hormone concentrations were similar in the two groups of patients. Diastolic relaxation was impaired to the same degree in the two groups of patients. Parameters of diastolic function showed no relation to left ventricular mass, which was significantly higher in the hypertensive than in the normotensive patients.

Conclusions—Uraemia is likely to induce specific changes in the relaxation properties of the myocardium. These changes are responsible for the impaired diastolic function independently of blood pressure, degree of hypertrophy, and metabolic changes, which suggests the existence of a specific cardiomyopathy. Hypertension remains a determinant of left ventricular mass.

Keywords: left ventricular morphology; diastolic function; uraemia; cardiomyopathy

Haemodialysis for chronic renal failure still carries high mortality (10% per year). Cardiovascular events are the main cause of death (50–60% against 15% in control populations).1 Accelerated coronary atherosclerosis, which has been claimed to be linked to lipid abnormalities, high blood pressure, diabetes, altered oxygen delivery at cellular level, and to an unknown uraemic factor, seems to be closely related to this excessive mortality.2 3

The natural course of disease in these patients is also characterised by congestive heart failure, possibly as a result of a specific uraemic cardiomyopathy.4 Several metabolic, ionic, hormonal, and biochemical factors have been suggested as the cause of this cardiomyopathy.5 6 Hyperparathyroidism may play an important part in the pathogenesis of ventricular dysfunction in uraemic patients, by determining inadequate left ventricular hypertrophy6 or by inducing low ventricular output and consequent failure and perhaps acting as a myocardial depressant.7 Recently, Horl and Riegel suggested that secondary hyperparathyroidism does not seem to be a clinically relevant myocardial depressant factor in uraemia, and they suggested the existence of a myocardial depressant factor of molecular weight between 10 000 and 30 000 daltons.8

The other factor invoked to explain ventricular hypertrophy and dysfunction is hypertension. In patients undergoing haemodialysis the prevalence of isolated systolic and systodias-astolic hypertension is fairly high. Simon et al reported a prevalence of 13% and 15% respectively.9 Parfrey and Harnett showed that systolic hypertension in uraemic patients is an independent risk factor for the development of left ventricular hypertrophy.10 In essential hypertension both systolic and diastolic high blood pressure determine left ventricular hypertrophy and eventually the systolic dysfunction that results in overt cardiac failure.11 However, the relation between hypertension, left ventricular hypertrophy, and diastolic function is not yet clear. In fact,
correlations between blood pressure measured in a doctor’s surgery and early alterations in diastolic function have not been found in essential hypertension, although a correlation was found with blood pressures monitored over 24 hours. Alterations in diastolic function are correlated with the progression of left ventricle hypertrophy, except in isolated systolic hypertension, where this correlation cannot always be found.

To verify whether a uraemic cardiomyopathy with specific myocardial features independent from hypertensive heart disease exists, we investigated the morphological profile and the diastolic function pattern of the left ventricle in a selected group of uraemic patients undergoing chronic haemodialysis by means of M mode, cross sectional, and Doppler echocardiography. A group of normal subjects with similar demographic characteristics was also studied.

Patients and methods

Sixty four patients receiving haemodialysis long term for chronic renal failure underwent echocardiographic and doppler assessment as screening for this project. They were all in sinus rhythm with neither physical nor laboratory signs of congestive heart failure. Medical history was negative for other disorders such as diabetes and haemochromatosis. Nineteen healthy normotensive subjects were chosen as controls from among the paramedical staff of this hospital; they were matched for age with the patients and had normal fasting glucose concentrations, lipid profiles, renal function, and packed cell volumes. Patients with pericardial effusion, constrictive pericarditis, haemodynamically significant valvar dysfunction, and regional dyskinesias were excluded. Thus the sample studied comprised only 35 (17 men, 18 women) of the 64 uraemic patients (54.7% of the sample).

All the patients had their blood tested; parathyroid hormone concentration was measured by means of a two site chemiluminescent immunometric assay of the entire molecule.

The presence of coronary heart disease was excluded by recalling the medical history and by means of electrocardiography and 24 hour Holter monitoring in which both the subcostal submaximal heart rate was reached. Echo measurements were carried out during the short interdiatilic interval (48 hours), exactly 24 hours after the end of the last dialysis. All the measurements were performed with a HP SONOS 1000, 2-5 and 3-5 mHz probes. The standardised method of Feigenbaum was used. Transmitral flow was assessed through an apical two or four chambers approach. Volume sample (4-5 mm length) was positioned within the mitral orifice at the level of the leaflets apex. Colour Doppler and signal frequency were used as a guide to align sample with the direction of flow.

With a parasternal short axis approach we measured left ventricular end diastolic and end systolic diameters, interventricular septum and posterior wall thicknesses, and systolic left atrium diameter. Aortic root diameter was measured with a parasternal long axis approach. Cardiac mass was measured by using the Devereaux equation. Ejection fraction was measured with the modified Simpson method in an apical two or four chamber approach. We also calculated the following Doppler parameters: maximal velocities of E and A waves (VmaxE, VmaxA); acceleration and deceleration of E and A waves (AccE, DecE; AccA, DecA); acceleration and deceleration times of E and A waves (DecTE, DecTA); the ratio of velocities in A and E waves (A/E); area of waves E and A (SE, SA); SA/SE + SA; and isovolumic relaxation time. Echo measurements were performed by a cardiologist and data analysed by two well trained observers; intrainsample and intersample variability were low (coefficient of variation 3%). Each measurement was done twice, and the average of the results obtained over three consecutive cardiac cycles was used.

Blood pressure was measured before and after each haemodialysis, but patients were allocated to the hypertensive or normotensive group by using the data from the 48 hours of continuous ambulatory monitoring, which was carried out in the short time between dialysis. Ambulatory monitoring data were also used because they correlate more strongly than blood pressure measured in a surgery with the prevalence of cardiovascular complications such as cardiac hypertrophy. A Spacelabs 90207 instrument was used for this purpose. Patients with a mean hourly blood pressure that was always lower than 160 or 95 mm Hg, or both, were considered to be normotensive. Patients with a mean hourly blood pressure greater than or equal to 160 or 95 mm Hg, or both, were considered to be hypertensive. Patients who had normal blood pressures after dialysis but pressures increased afterwards to mean hourly values greater than 160 or 95 mm Hg, or both, were considered to have volume dependent hypertension and were included in the hypertensive group.

Statistical analysis was carried out by means of a BMDF statistical package. One way analysis of variance was used to compare the three groups of parameters. Multiple analysis of covariance was used for the indices of cardiac hypertrophy in the two subgroups of uraemic patients. Within the uraemic patients linear regression analysis was used to correlate cardiac mass with the mean systolic and diastolic blood pressures over 48 hours and with the diurnal and nocturnal values; cardiac mass with indices of diastolic function; ambulatory blood pressures with parameters of diastolic function; parathyroid hormone concentrations with indices of diastolic function and ambulatory blood pressure over 48 hours. A two way analysis of variance was carried out in uraemic patients to see whether differences in cardiac mass were related to sex.
Table 1 Clinical and laboratory data on uraemic patients. Values are mean (SD)

<table>
<thead>
<tr>
<th></th>
<th>All (17 men, 18 women)</th>
<th>Normotensive (5 men, 6 women)</th>
<th>Hypertensive (12 men, 10 women)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>60 (14)</td>
<td>60 (14)</td>
<td>59 (16)</td>
</tr>
<tr>
<td>Duration of dialysis (months)</td>
<td>52 (54)</td>
<td>53 (40)</td>
<td>40 (57)</td>
</tr>
<tr>
<td>Measurements before dialysis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haemoglobin (g/l)</td>
<td>85 (4)</td>
<td>85 (3)</td>
<td>86 (4)</td>
</tr>
<tr>
<td>Packed cell volume (%)</td>
<td>30 (4-4)</td>
<td>30 (5-4)</td>
<td>30 (8-7)</td>
</tr>
<tr>
<td>Sodium (mmEq)</td>
<td>139 (1-3)</td>
<td>138 (1-3)</td>
<td>139 (1-1)</td>
</tr>
<tr>
<td>Potassium (mmEq)</td>
<td>5.2 (0.5)</td>
<td>5.3 (0.4)</td>
<td>5.1 (0.5)</td>
</tr>
<tr>
<td>Calcium (mg/l)</td>
<td>94 (20)</td>
<td>93 (21)</td>
<td>95 (98)</td>
</tr>
<tr>
<td>Blood urea nitrogen (mg/l)</td>
<td>846 (167)</td>
<td>847 (167)</td>
<td>855 (122)</td>
</tr>
<tr>
<td>Phosphataemia (g/l)</td>
<td>43 (4)</td>
<td>43 (4)</td>
<td>43 (3)</td>
</tr>
<tr>
<td>Alkaline phosphatase (IU/l)</td>
<td>106 (50-6)</td>
<td>106 (50-6)</td>
<td>106 (50-6)</td>
</tr>
</tbody>
</table>

Table 2 Mean (SD) blood pressures (mm Hg) in controls and uraemic patients

<table>
<thead>
<tr>
<th></th>
<th>Controls (6 men, 13 women)</th>
<th>Normotensive (5 men, 8 women)</th>
<th>Hypertensive (12 men, 10 women)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systolic</td>
<td>113.8 (5.4)*</td>
<td>110.4 (13.5)</td>
<td>145.1 (13.5)*</td>
</tr>
<tr>
<td>Diastolic</td>
<td>67.9 (4.6)*</td>
<td>64.0 (8.5)</td>
<td>86.7 (11.1)*</td>
</tr>
<tr>
<td>Daytime:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Systolic</td>
<td>115.8 (6.4)*</td>
<td>111.7 (13-0)</td>
<td>145.9 (14.5)*</td>
</tr>
<tr>
<td>Diastolic</td>
<td>69.9 (5.8)*</td>
<td>65.5 (8-1)</td>
<td>87.4 (11.4)*</td>
</tr>
<tr>
<td>Night time:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Systolic</td>
<td>110.5 (6.6)†</td>
<td>106.2 (14.1)</td>
<td>144.6 (15.8)*</td>
</tr>
<tr>
<td>Diastolic</td>
<td>65.1 (4.7)†</td>
<td>61.6 (9-1)</td>
<td>86.0 (15-1)*</td>
</tr>
</tbody>
</table>

*P < 0.001 vs uraemic normotensive patients. **P < 0.01 vs uraemic normotensive patients. †P = NS vs uraemic normotensive patients. §P < 0.001 vs controls.

Results

Table 1 shows the baseline characteristics of the uraemic patients. Their mean age was 60-3 (14-2) years and they had received dialysis for a mean of 52 months (range 10-76). 66% of the patients were treated with h-erythropoetin to achieve a target haemoglobin concentration of 8-5 g/l (packed cell volume 30%). According to the results of blood pressure monitoring over 48 hours, 22 patients (12 men, 10 women) were categorised as being hypertensive and 13 patients (five men, eight women) as normotensive. Controls (six men and 13 women) had a mean age of 57-5 (8-4) years.

The original nephropathy in the 35 uraemic patients was undetermined nephropathy (eight patients), nephroangiosclerosis (nine), rapidly progressive glomerulonephritis (two), interstitial nephropathy (five), obstructive nephropathy (two), chronic glomerulonephritis (six), and adult polycystic kidney disease (three) (P2 = 3-61, NS).

Table 2 shows the ambulatory blood pressures over 48 hours in the three groups. Control subjects had similar blood pressures to normotensive uraemic patients, but significantly lower blood pressures than the hypertensive uraemic patients (P < 0.001 for all the variables analysed). Systolic and diastolic blood pressures were always significantly higher in the hypertensive patients than in the normotensive patients.

Multiple logistic regression analysis with hypertension as the dependent variable was not related to sex, blood urea nitrogen, creatinine concentration, phosphataemia, serum albumin concentration, parathyroid hormone concentration, serum concentrations of calcium, sodium, and potassium, packed cell volume, or KTV as predictors. Parathyroid hormone concentration was 229.9 (276-0) nmol/l in the normotensive uraemic patients and 247-9 (207-4) nmol/l in the hypertensive uraemic patients (normal range 5–50 nmol/l).

Table 3 summarises the echocardiographic measurements in the three groups. Interventricular septal and posterior wall thicknesses, systolic left atrial diameter, and aortic root diameter were significantly higher in the uraemic patients (P < 0.0001). Left ventricular end diastolic diameter and cardiac mass were also significantly different (P < 0.001 and P = 0.017 respectively).

No significant differences were found in ejection fraction. When the two subgroups of hypertensives and normotensives were compared, only the interventricular septal and posterior wall thicknesses and cardiac mass were significantly higher in the first group. There was no overlap between uraemic and control patients in interventricular septal and posterior wall thicknesses. Overlap was almost completely absent between hypertensive and normotensive patients undergoing dialysis. Table 4 shows the various parameters of left ventricular diastolic function. VmaxA, DcA, AccA, DecTE, DecTA, A/E, SA, S/SA + SE, and isovolumic relaxation time were significantly higher in uraemic patients than controls.

A multiple analysis of covariance was carried out to see whether differences in diastolic function between hypertensive and normotensive patients undergoing dialysis could be accounted for by the different degree of hypertrophy. Diastolic function parameters were therefore analysed with left ventricular mass index, and posterior wall and interventricular septal thicknesses used as covariates. The two subgroups of patients did not differ in VmaxA, DcE, AccE, DecTE, DecTA, A/E, SA, S/SA + SE, and isovolumic relaxation time were significantly higher in uraemic patients than controls.

There was a significantly positive correla-
and the alterations in diastolic function. A cut off point between hypertension and normotension in uraemic patients of 160/90 mm Hg might seem arbitrary, but this was an attempt to include isolated systolic hypertension (defined in large clinical trials such as SHEP and SYST-EUR greater than 160 mm Hg) and systodiastolic hypertension. A similar cut off point was adopted in a previous study using ambulatory blood pressure monitoring in patients undergoing haemodialysis.

Our morphological data are similar to published data; in fact, uraemic patients have much thicker ventricle walls, a greater left ventricular mass, and a higher degree of dilatation. Within these patients, hypertensive subjects show a significantly higher left ventricular mass and thicker interventricular septum. However, the presence of morphological changes even in the subgroup of normotensive uraemic patients indicates that chronic renal failure or its inadequate correction by haemodialysis may have an important and independent role. Hypertension can worsen the degree of hypertrophy. Left ventricular mass correlates significantly with the average diastolic blood pressure and the mean systolic and diastolic nocturnal pressures over 48 hours, which can be considered to be good indices of cardiac workload. Patients considered normotensive in this study might have been hypertensive in the past and have therefore developed some degree of cardiac hypertrophy. If that had been the case, lowering blood pressure should none the less have produced regression in left ventricular hypertrophy. In terms of systolic dysfunction in our uraemic patients, this is present to a lesser degree than reported in other series, but this can be explained, at least in part, by the fact that we excluded all the subjects with haemodynamically significant valve dysfunction, ischaemia, and supraventricular tachyarrhythmias such as atrial fibrillation. Recently the importance of studying not only the anatomical but also the functional diastolic changes in the left ventricle have been emphasised. This can be easily done by means of Doppler echocardiography of transmural flow, such data being consistent with the data obtained by other methods such as angiography and radionuclide scanning.

Determinants of transmural flow are the active relaxation of the left ventricle, myocardial stiffness, and some haemodynamic variables such as preload and afterload, atrial pressure, and ventricular pressure. Pressure and volume overload are characterised by typical modifications of the diastolic curves. Patients undergoing dialysis long term show an altered diastolic function similar to that of patients with left ventricular hypertrophy without interdialytic volume overload, though they may present with a pressure and volume overload. Transmural flow is characterised by a dominant A wave with an increased isovolumic relaxation time. In clinical and experimental models the increased ventricle stiffness which accompanies left ventricular hypertrophy is mainly due to changes in the collagen

### Table 4: Mean (SE) transmural flow Doppler measurements

<table>
<thead>
<tr>
<th>Patients with uraemia</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Normotensive</td>
</tr>
<tr>
<td>VmaxE (cm/s)</td>
<td>69.9 (3.2)</td>
</tr>
<tr>
<td>AccE (cm/s)</td>
<td>896.3 (40.2)</td>
</tr>
<tr>
<td>DecE (cm/s)</td>
<td>527.7 (50.9)*</td>
</tr>
<tr>
<td>AccTE (m/s)</td>
<td>79.0 (7.2)</td>
</tr>
<tr>
<td>DecTE (m/s)</td>
<td>131.1 (8.8)</td>
</tr>
<tr>
<td>SE (cm)</td>
<td>10.3 (0.7)</td>
</tr>
<tr>
<td>VmaxA (cm/s)</td>
<td>64.7 (3.8)*</td>
</tr>
<tr>
<td>AccA (cm/s)</td>
<td>887.0 (70.8)*</td>
</tr>
<tr>
<td>DecA (cm/s)</td>
<td>630.0 (56.7)</td>
</tr>
<tr>
<td>AccTA (m/s)</td>
<td>60.7 (5.9)</td>
</tr>
<tr>
<td>DecTA (m/s)</td>
<td>79.5 (6.8)*</td>
</tr>
<tr>
<td>SA (cm)</td>
<td>6.4 (0.5)</td>
</tr>
<tr>
<td>ISA</td>
<td>0.40 (0.02)*</td>
</tr>
<tr>
<td>A/E (cm/s)</td>
<td>1.0 (0.1)*</td>
</tr>
<tr>
<td>IVRT (m/s)</td>
<td>76.0 (5.7)*</td>
</tr>
</tbody>
</table>

One way analysis of variance was used.

VmaxE, E wave maximal velocity; AccE, E wave acceleration; DecE, E wave deceleration; AccTE, E wave acceleration time; DecTE, E wave deceleration time; SE, E wave area; VmaxA, A wave maximal velocity; AccA, A wave acceleration; DecA, A wave deceleration; AccTA, A wave acceleration time; DecTA, A wave deceleration time; SA, A wave area; ISA, SA/SA + SE; A/E, A wave area/E wave area; IVRT, isovolumic relaxation time.

*Significantly different patients with uraemia.

### Discussion

Chronic renal failure is accompanied by a high prevalence of morphological and functional changes in the heart, and these can easily be detected by echocardiography. These abnormalities are responsible for the high incidence of congestive heart failure and cardiac mortality in affected patients. Many authors ascribed these changes to coronary heart disease or hypertension, which are both often found in uraemic patients receiving dialysis. These patients might develop a specific cardiomyopathy, secondary to their specific metabolic, biochemical, hormonal, and maybe even haemodynamic features (atrioventricular fistula). In the early phases of uraemic cardiomyopathy the interventricular septum and the posterior wall become thickened in association with appreciable dilatation of the left atrium in 40% of patients. Global dilatation of the left ventricle occurs only in the late stages, and some authors consider this to be the result of high blood pressure, which is often found in these subjects, rather than of intrinsic changes due to uraemia.

Our blood pressure data are from 48 hour monitoring; in other studies such data seem to be closely correlated to the prevalence of complications, the degree of cardiac hypertrophy, and the alterations in diastolic function. A cut off point between hypertension and normotension in uraemic patients of 160/90 mm Hg might seem arbitrary, but this was an attempt to include isolated systolic hypertension (defined in large clinical trials such as SHEP and SYST-EUR greater than 160 mm Hg) and systodiastolic hypertension. A similar cut off point was adopted in a previous study using ambulatory blood pressure monitoring in patients undergoing haemodialysis.

Our morphological data are similar to published data; in fact, uraemic patients have much thicker ventricle walls, a greater left ventricular mass, and a higher degree of dilatation. Within these patients, hypertensive subjects show a significantly higher left ventricular mass and thicker interventricular septum. However, the presence of morphological changes even in the subgroup of normotensive uraemic patients indicates that chronic renal failure or its inadequate correction by haemodialysis may have an important and independent role. Hypertension can worsen the degree of hypertrophy. Left ventricular mass correlates significantly with the average diastolic blood pressure and the mean systolic and diastolic nocturnal pressures over 48 hours, which can be considered to be good indices of cardiac workload. Patients considered normotensive in this study might have been hypertensive in the past and have therefore developed some degree of cardiac hypertrophy. If that had been the case, lowering blood pressure should none the less have produced regression in left ventricular hypertrophy. In terms of systolic dysfunction in our uraemic patients, this is present to a lesser degree than reported in other series, but this can be explained, at least in part, by the fact that we excluded all the subjects with haemodynamically significant valve dysfunction, ischaemia, and supraventricular tachyarrhythmias such as atrial fibrillation. Recently the importance of studying not only the anatomical but also the functional diastolic changes in the left ventricle have been emphasised. This can be easily done by means of Doppler echocardiography of transmural flow, such data being consistent with the data obtained by other methods such as angiography and radionuclide scanning.

Determinants of transmural flow are the active relaxation of the left ventricle, myocardial stiffness, and some haemodynamic variables such as preload and afterload, atrial pressure, and ventricular pressure. Pressure and volume overload are characterised by typical modifications of the diastolic curve. Patients undergoing dialysis long term show an altered diastolic function similar to that of patients with left ventricular hypertrophy without interdialytic volume overload, though they may present with a pressure and volume overload. Transmural flow is characterised by a dominant A wave with an increased isovolumic relaxation time. In clinical and experimental models the increased ventricle stiffness which accompanies left ventricular hypertrophy is mainly due to changes in the collagen-
network rather than to the increased myocyte mass. Recent data in uraemic patients and rats suggest that uraemia is characterised by an increased myocardial collagen content in the myocardium—so called diffuse intermyocardiotic fibrosis—although myocyte hypertrophy has l

do been described. The altered diastolic function seems therefore to be linked to fibrosis. Lowering blood pressure by means of methylodopa, calcium channel blockers, angiotensin converting enzyme inhibitors, and canrenone potassium 42-44 in experimental models of left ventricular hypertrophy decreases left ventricular mass by reducing myocyte hypertrophy. Collagen content is also decreased and collagen phenotypes reshifted.

We therefore suggest that the altered diastolic function in our patients can be mainly ascribed to an increased myocardial stiffness since changes in the A wave and isovolumic relaxation time are more pronounced. Changes in myocardial function due to increased preload have a minor role. Our hypertensive and normotensive patients showed the same degree of changes in diastolic function, though they had significantly different indices of left ventricular mass and none of the diastolic parameters correlated with left ventricular mass or with blood pressures over 48 hours. These data are only partially in keeping with those reported for uraemic patients. Huting et al reported that diastolic function does not correlate with the degree of left ventricular hypertrophy, patients with uraemia showing impaired diastolic filling independently from left ventricular mass. However, the same group found a correlation between diastolic function and blood pressure measured in a doctor's surgery. Myocardial stiffness in uraemic patients seems to be complex, with many factors contributing to the development of cardiac hypertrophy. This is not the case for left ventricular mass, which correlates well with blood pressure monitored over 48 hours and therefore with cardiac workload.

We think that the increased myocardial stiffness is determined also by uraemia. In fact, we recently showed that contractility of single isolated myocytes, as well as their intrinsic speed of shortening and relaxation, is depressed in rats with chronic uraemia. These changes were not ascribed to myocyte hypertrophy since contractile proteins did not show the typical shift towards the forms of the hypertrophied myocardium. We hypothesised that cardiotocic substances produced during uraemia could slow down contraction and relaxation.

Hyperparathyroidism may have an important role in the pathogenesis of uraemic cardiomyopathy. Indeed, our patients have high concentrations of parathyroid hormone regardless of their blood pressure. We found no correlations between parathyroid hormone concentration and indices of diastolic function, but we cannot say that abnormal diastolic function is not affected by parathyroid hormone. The same may be true for other variables such as haemoglobin, ions, and other metabolites, whose concentrations were almost identical in the two groups of uraemic patients. Recently, however, “uraemic cardiodiodepressant factors” with molecular weights between 10 000 and 30 000 daltons have been described which may modify the composition of the interstitium, as well as metabolic factors, ions, and hormones. Further studies are needed to identify a cause and effect relation.


27. Torok E, Borbas S, Lengyel M, Zorandi A: Regression of cardiac hypertrophy in hypertensive patients by long-
Left ventricular morphology and diastolic function in uraemia: echocardiographic evidence of a specific cardiomyopathy.


Br Heart J 1995 74: 174-179
doi: 10.1136/hrt.74.2.174