Familial and primary (AL) cardiac amyloidosis: echocardiographically similar diseases with distinctly different clinical outcomes

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Abstract

Objective—To determine whether patients with myocardial amyloidosis due either to AL (primary) amyloid or familial amyloid have distinguishing echocardiographic or electrocardiographic features; and to compare the prevalence of heart failure and survival in the two types of amyloidosis in relation to echocardiographic findings.

Design—Blinded group comparison of randomly selected cases of cardiac amyloidosis.

Setting—International referral centre for amyloid research and treatment.

Patients—36 patients with cardiac amyloid heart disease, of whom 12 had familial and 24 had primary AL amyloidosis.

Results—Familial and AL echocardiograms were morphologically indistinguishable, with similar left ventricular wall thickness, mean (SD) 15-4 (2-3) v 15-8 (2-5) mm, respectively; right ventricular wall thickness was also similar between amyloid types: 9-6 (2-8) v 9-7 (6-5) mm, respectively. Doppler indices of left and right ventricular function, left ventricular volume, and ejection fraction were also similar. Low voltage electrocardiograms (<0-5 mV) were more common in the AL (16/24, 67%) than in the familial group (4/12, 25%), P < 0-05. The one year survival for familial and AL forms was 92% (11/12) v 38% (6/24), respectively, with virtually all deaths due to cardiac causes.

Conclusions—Although cardiac involvement is echocardiographically indistinguishable, cardiac mortality is very different between the two forms of amyloidosis. Preservation of echocardiographic voltage in familial amyloidosis suggests that the particular biochemical characteristics of distinct types of amyloid fibril have different pathological effects on the myocardium. This distinction becomes critical in the evaluation, treatment, and management of patients who have a diagnosis within the spectrum of the protein deposition diseases.

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Keywords: amyloidosis; echocardiography; familial amyloidosis

Familial amyloid polyneuropathy (FAP), identified just over 40 years ago, is the rarest type of systemic amyloidosis. The exact prevalence of FAP is unknown but, based on referral patterns to major centres specialising in amyloidosis, it probably represents 10% of patients with the disease. It is estimated that there are between 1000 and 2225 new cases of AL (primary) amyloidosis in the United States each year, and therefore one might predict between 100 to 200 newly diagnosed cases of familial amyloidosis per year. A transthyretin mutant protein produced by the liver is responsible for most cases of familial amyloidosis, and the resultant protein deposition may cause dysfunction of various organ systems including the peripheral and autonomic nerves, gastrointestinal tract, and heart. Diagnosis of conduction necessitating permanent pacing are the predominant manifestation of cardiovascular disease, but myocardial infiltration with amyloid is also frequently present.

The echocardiographic appearance of myocardial involvement in FAP has been described as similar to that in AL amyloidosis (light chain amyloidosis, formerly called primary amyloidosis). Typical abnormalities consist of increased right and left ventricular wall thickness with normal cavity size, increased myocardial echogenicity, and valve thickening. Data from patients with AL amyloid indicate that myocardial involvement documented by echocardiography is often accompanied by congestive heart failure and that its presence is associated with a median survival less than six months. Survival in familial amyloidosis with cardiac involvement appears to be longer than in AL amyloidosis, although no direct comparison has been made and no attempt has been made to compare groups of patients with similar echocardiographic appearances. Recently liver transplantation has been shown to be an effective treatment for patients with FAP. By removing the source of the mutant protein the disease is arrested and may even regress. The significance of cardiac amyloid infiltration in patients being evaluated for liver transplantation is not known. However, since its presence in AL amyloidosis augurs an ominous prognosis, it might be expected to carry a similar prognosis in the familial form, thus rendering the risks of major surgery prohibitive. We therefore undertook this study to determine whether patients with FAP and definite cardiac involvement had echocardiographic features which distinguished them from patients with AL cardiac amyloidosis, and whether the clinical and prognostic significance of such cardiac involvement differed from a cohort of patients with AL amyloidosis.
Methods

Over a period of eight years (1986–94), 40 patients with familial amyloidosis and 232 with AL amyloidosis were seen at the clinical research centre of Boston University Medical Center. All patients referred with the diagnosis of amyloidosis underwent an extensive clinical evaluation including a full non-invasive cardiovascular examination. This includes a 12 lead electrocardiogram, 24 hour Holter recording, and Doppler and cross sectional echocardiography. Following initial evaluation patients are prospectively followed by written contact and annual visits.

Echocardiography

Echocardiograms were performed using a Hewlett Packard phased array system. M mode recordings were made at 50 mm/s with simultaneous recording of the electrocardiogram. Heart failure was considered present on physical examination in patients with raised jugular venous pressure or evidence of pulmonary venous congestion or both. Severity of heart failure was classified using the New York Heart Association criteria.22 Of these 40 patients with FAP, 17 had no clinical or echocardiographic evidence of cardiac involvement and two had conduction system disease but normal echocardiograms. An additional nine patients had mild left ventricular wall thickening on echo (≤ 1.3 cm) without clinical symptoms of cardiac disease. The remaining 12 patients from the familial group had ventricular wall thickening greater than 1.3 cm unexplained by hypertension or valve disease, and formed the index group for this study. In order to evaluate similarities and differences between cardiac involvement in patients with FAP and AL amyloidosis we randomly chose 24 patients with AL amyloidosis from among 133 patients whose echocardiograms showed a wall thickness > 1.3 cm, in the absence of hypertension or significant valve disease. Cardiac involvement by echocardiography was the single criterion for patient recruitment of both groups and selection was blind to any additional clinical features or outcome aside from amyloid type.

Diagnosis of Amyloid Type

All 36 patients studied had biopsy proven systemic amyloidosis. The diagnosis of AL amyloidosis was made by histological evidence of systemic amyloid in association with evidence of a plasma cell dyscrasia and/or by identification of an immunoglobulin light chain in their serum, urine, or amyloid deposit. None of these 24 patients had evidence of multiple myeloma. The diagnosis of FAP was confirmed by isoelectric focusing of the serum to identify variant transthyretin, which was followed by detection of a mutant transthyretin gene known to be associated with familial amyloidosis identified in their DNA. In all 12, a family history of amyloidosis was also present. The type of transthyretin mutation responsible for amyloid production was available in 37 of the 40 FAP patients (92.5%) from whom the study group was derived. The commonest mutation, occurring in 10 of the 40 patients (25%), was valine-30-methionine. This mutation was only present in two of the 12 FAP patients (16.7%) with significant echocardiographic abnormalities, reflecting the relative infrequency of myocardial infiltration in this genetic form compared with other mutations. The remaining 10 FAP index group patients had the following amino acid substitutions: threonine-60-alanine in two, valine-30-alanine in one, and glutamate-89-glutamine in one. Six further patients were identified as having two mutations each, respectively glutamate-42-glycine and histidine-90-asparagine.

Survival Analysis

Survival was measured from the time of the echocardiogram until death except in four AL patients who underwent cardiac transplantation. These patients were considered to have fatal organ failure (a death equivalent) from the date of this procedure. Liver transplantation was not considered an end point in patients with FAP, since the liver function in these patients is normal and the purpose of transplantation is to remove the chronic source of mutant protein synthesis. Within the AL group, 11 patients received melphalan and prednisone, 12 received colchicine alone, and one patient had an extremely rapid demise and died before any treatment could be started.

AL amyloid may seriously affect the kidneys as well as the heart and this combination may have a detrimental effect on survival. We addressed any unintentional bias towards more significant renal disease in the AL group by reanalysing the AL patients (n = 16) without such renal disease (serum creatinine < 115 μmol/l and 24 hour urinary protein excretion < 3 g/24 hours on any occasion) and comparing the survival of these 16 AL patients with the familial group. We also compared survival in a subgroup of the AL patients with cardiac amyloidosis who were age matched to the familial group.

Assessment of Echocardiograms

This was performed for all 36 echocardiograms, in a blinded fashion, to determine if there were features that were specific to the individual types of amyloidosis. Two experienced cardiologists examined the echocardiograms; both were unaware of the type of amyloid disease. For all patients a consensus opinion was reached regarding the presence of the classical features originally described in AL cardiac amyloidosis7: myocardial thickening and increased echogenicity ("granular sparkling"), valve thickening, and interatrial septal thickening. Additional features previously described in amyloid heart disease, including pericardial effusion and atrial dilatation, were also recorded. Left ventricular systolic performance was judged to be normal (ejection fraction ≥ 55%), mildly impaired (ejection fraction of 40–54%), moderately impaired (ejection fraction 30–39%), or severely impaired (ejection fraction < 30%).

In addition to blinded qualitative assess-
Figure 1. Illustration of Doppler measurements and intervals. Aortic flow is represented as the waveform below the baseline. AC represents aortic valve closure; DT, the deceleration time of the early filling phase; S1, the first heart sound; IT, the left ventricular isovolumetric relaxation time, measured from the aortic valve closure (AC) to the start of mitral flow (MO). The shaded areas of the mitral waveform indicate the waveform proportions used in the measurement of the E and A wave velocity time integrals.

Doppler, Cha, Skinner, LaValley, Falk

ELECTROCARDIOGRAPHY

Twelve lead electrocardiograms were performed on all patients and calibrated to a 10 mm deflection equivalence to 1 mV. A low voltage electrocardiogram was defined as a mean QRS voltage amplitude in the standard and unipolar leads (I, II, III, aVL, and aVF) of \( \leq 0.5 \) mV. A pseudoinfarction pattern was defined by QS waves in the anteroseptal (V1-V3) and/or the inferior leads (II, III, and aVF), in the absence of previous myocardial infarction. Augmented left ventricular voltages were said to be present if the S wave in V1 plus the largest R wave in V5 or V6 exceeded 3.5 mV. The duration of the QT interval was corrected for heart rate to produce the Qtc interval.17 Prolongation of the Qtc interval was taken as a duration of more than 0.45 seconds for men and 0.47 seconds for women.18 Twenty four hour Holter recordings were analysed for the presence of rhythm disturbances and conduction disorders.

Electrocardiographic voltage to left ventricular mass ratio

The voltage to mass ratio was derived from the electrocardiogram voltage both for the precordial leads (S wave voltage in V1 plus the mean of the R wave voltage in V5 and V6),19 and the limb leads. This was divided by the cross sectional area of the left ventricular wall (CSA) imaged in a transverse plane.20 CSA was calculated from the left ventricular end diastolic dimension (LVEDD), the mean left ventricular wall thickness (Th) at end diastole, and the body surface area (BSA), using the geometric formula:

\[
CSA = \left[ \frac{(LVEDD/2 + Th)^2 - (LVEDD/2)^2}{2} \right] / BSA
\]

In healthy subjects without cardiac amyloidosis normal values for this left ventricular mass equivalent corrected for body surface area are in the range of 6–10 cm²/m².20

STATISTICAL ANALYSIS

Values are given as mean (SD). Statistical analyses were carried out using the SAS package (SAS Institute, Cary, North Carolina, USA). Continuous data for Doppler, echocardiographic, and electrocardiographic measurements were tested using an unpaired non-parametric Mann Whitney U test. Fisher's exact test was used for the analysis of discrete
Table 1 Clinical and laboratory features of the two patient groups with cardiac amyloidosis

<table>
<thead>
<tr>
<th>Features</th>
<th>Familial (FAP) amyloidosis (n = 12)</th>
<th>Primary (AL) amyloidosis (n = 24)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>47 (11)</td>
<td>58 (11)</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Sex</td>
<td>M/F</td>
<td>M/F</td>
<td></td>
</tr>
<tr>
<td>Systolic BP (mm Hg)</td>
<td>114 (11)</td>
<td>113 (11)</td>
<td>NS</td>
</tr>
<tr>
<td>Diastolic BP (mm Hg)</td>
<td>72 (8)</td>
<td>68 (11)</td>
<td>NS</td>
</tr>
<tr>
<td>Heart rate (beats/min)</td>
<td>83 (15)</td>
<td>75 (6)</td>
<td>NS</td>
</tr>
<tr>
<td>Total serum protein (g/dl)</td>
<td>6-1 (0-7)</td>
<td>5-8 (0-8)</td>
<td>NS</td>
</tr>
<tr>
<td>Serum albumin (g/dl)</td>
<td>3-7 (0-5)</td>
<td>3-5 (0-6)</td>
<td>NS</td>
</tr>
</tbody>
</table>

Data are reported as mean (SD) or No (%). Median survival is measured from date of the echocardiogram.

BP, blood pressure; CHF, congestive heart failure; NYHA, New York Heart Association classification of heart failure.

* = n = 23 for primary group as one patient in this group had intermittent symptoms of CHF and angina that was also influenced by concurrent renal dialysis, thus making the classification of NYHA heart failure unreliable in this patient.

Results

Twelve patients (eight men, four women) formed the familial group, with a mean age of 47 (11) years. The AL amyloid group consisted of 24 patients (15 men, nine women) with a mean age of 58 (11) years; significantly greater than that of the familial group (P < 0.01). All patients were in sinus rhythm. The heart rate and both systolic and diastolic blood pressures were similar for patients with the two types of amyloidosis (table 1). Significant multigorgan involvement was present in eight of 12 FAP patients; six had gastrointestinal and neurological symptoms, while two had gastrointestinal and four had neurological symptoms in addition to their cardiac disease. Renal involvement, with a serum creatinine > 176 μmol/l and/or a 24 hour urine protein > 3 g, was present in eight of the 24 patients with AL amyloidosis. However serum albumin was not below 2 g/dl in any patient.

Echocardiographic Differences Between FAP and AL Amyloid Groups

On blinded review of the echocardiograms, the readers were unable to distinguish AL amyloidosis from FAP (fig 2) and no differences in echocardiographic features were noted between groups (table 2). In addition to the well described features of cardiac amyloidosis, that is, increased myocardial echogenicity, and thickening of walls, valves, and the atrial septum, we found that pericardial effusion and atrioventricular valve incompetence was common. Atrioventricular valve incompetence occurred in 92% (11 of 12) of the familial and 88% (21 of 24) of the AL amyloid groups, with a trend for more of the AL amyloid patient group to have higher grades of tricuspid incompetence (table 2).

Quantitative echocardiographic measurements

Mean values for left atrial diameter and interventricular septal, left ventricular posterior...
between the two amyloid groups for the derived ejection fraction (table 3).

**Doppler evaluation of ventricular inflow, mitral E wave deceleration time and left ventricular isovolumetric relaxation time**

Doppler evaluation of mitral valve flow failed to show any differences between groups in A wave velocity, E wave velocity, E to A peak velocity ratio, or ratio of E to A velocity time integrals. Although the mean E wave and A wave velocities for the two groups each fell within the normal range, several individuals in each group had very small A waves, and two patients in the AL amyloid group had absent A wave flow on Doppler examination. As a result, mean E to A ratios for both groups lay well above the normal range (table 4).  

Mitral E wave flow deceleration time was similarly reduced in both the familial (157 (28) ms) and AL amyloid groups (151 (28) ms) when compared to normal reference values. The left ventricular isovolumetric relaxation time was similar in patients from the FAP (89 (18) ms) and the AL patient groups (92 (25) ms). Criteria for a restrictive left ventricular filling pattern on Doppler were satisfied by three of the 12 FAP patients (25%) and by 12 of the 24 AL patients (50%).

Peak tricuspid flow velocity and flow velocity integral in early diastole (E wave flow) were similar in both the FAP and AL groups. Similarly, the peak tricuspid flow velocity and flow velocity integral with atrial contraction in late diastole (A wave flow) did not differ between the two groups. The resultant E to A flow velocity ratio was similar in both the FAP (1·26 (0·46)) and AL groups (1·98 (1·34)).

The flow velocity integral of the aortic outflow Doppler was greater in the FAP patients (17·78 (5·34) cm) than in the AL patients (14·28 (4·87) cm, P < 0·05), although calculated cardiac output did not achieve statistical significance (table 4).

**ELECTROCARDIOGRAPHIC DATA**

Common abnormal features included pseudo-infarction and abnormal axis deviation (table 5). Of note, no patient in either group had left bundle branch block. Mean limb lead voltage was lower in the AL patients (0·42 (0·20) mV) than in the FAP group (0·68 (0·29) mV, P = 0·01) (figs 3 and 4). When expressed as a left ventricular voltage (limb leads) to mass equivalent ratio there remained a significant difference between the FAP (0·21 (0·11)) and AL groups (0·14 (0·06), P < 0·05).

**TWENTY FOUR HOUR HOLTER DATA**

Both patient groups showed similar numbers of atrial and ventricular premature beats, and episodes of ventricular and supraventricular tachycardia, including atrial fibrillation and flutter (table 5). Two patients in the AL group had had permanent pacemaker implantation; of these one was a patient with second degree atrioventricular block and the second was in a patient who had documented electromechanical dissociation and cardiac arrest during the early stages of a modified exercise stress test.
Table 5 Values for resting 12 lead and 24 hour Holter electrocardiographic recordings in the two patient groups with cardiac amyloidosis

<table>
<thead>
<tr>
<th>Feature</th>
<th>Familial (FAP) amyloidosis (n = 12)</th>
<th>Primary (AL) amyloidosis (n = 24)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abnormal ECG</td>
<td>11 (92%)</td>
<td>22 (92%)</td>
<td>NS</td>
</tr>
<tr>
<td>Low voltage ECG</td>
<td>4 (33%)</td>
<td>10 (67%)</td>
<td>NS</td>
</tr>
<tr>
<td>Axis deviation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right: (+90° to +180°)</td>
<td>4 (33%)</td>
<td>8 (33%)</td>
<td>NS</td>
</tr>
<tr>
<td>Left: (-30° to -90°)</td>
<td>4 (33%)</td>
<td>8 (33%)</td>
<td>NS</td>
</tr>
<tr>
<td>ECG voltage (limb) (mV)</td>
<td>0.68 (0.29)</td>
<td>0.42 (0.20)</td>
<td>0.01</td>
</tr>
<tr>
<td>LV-CSA/BSA (cm²/m²)</td>
<td>17.2 (3.8)</td>
<td>16.0 (2.9)</td>
<td>NS</td>
</tr>
<tr>
<td>ECG voltage (limb): LVMR</td>
<td>0.21 (0.10)</td>
<td>0.14 (0.06)</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Pseudoinfarct pattern – inferior</td>
<td>3 (25%)</td>
<td>6 (25%)</td>
<td>NS</td>
</tr>
<tr>
<td>Pseudoinfarct pattern – anterior</td>
<td>6 (50%)</td>
<td>9 (38%)</td>
<td>NS</td>
</tr>
<tr>
<td>Conduction disturbances</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1st degree AV block</td>
<td>3 (25%)</td>
<td>4 (17%)</td>
<td>NS</td>
</tr>
<tr>
<td>2nd degree AV block</td>
<td>1 (4%)</td>
<td>2 (8%)</td>
<td>NS</td>
</tr>
<tr>
<td>Bifascicular block</td>
<td>3 (25%)</td>
<td>2 (8%)</td>
<td>NS</td>
</tr>
<tr>
<td>24 Hour Holter data</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VT</td>
<td>1 (8%)</td>
<td>1 (4%)</td>
<td>NS</td>
</tr>
<tr>
<td>SVT</td>
<td>1 (8%)</td>
<td>3 (13%)</td>
<td>NS</td>
</tr>
<tr>
<td>Atrial fibrillation or flutter</td>
<td>2 (17%)</td>
<td>1 (4%)</td>
<td>NS</td>
</tr>
</tbody>
</table>

Data are reported as mean (SD) or No (%).

In the FAP group two patients also required permanent pacemakers, the indications in both cases being symptomatic bradycardia with syncope.

**SURVIVAL CHARACTERISTICS**

Patients in the FAP group had a 92% (11 of 12) one year survival from the date of their echocardiogram, compared to 25% (six of 24) for those patients in the AL group (P = 0.0002) (table 1). Reference to the Kaplan-Meier survival analysis (fig 5) shows that prolonged survival was frequent in the FAP patients and not in the AL patients (P < 0.0002). One AL patient died of hepatic failure and the remaining 22 deaths in this group were cardiac: sudden in six and from congestive heart failure in 16. Of the five deaths in the FAP patients, one was caused by sepsis, one was due to inanition, and three were possibly cardiac—two being sudden and one followed several hours of breathlessness of unknown aetiology (table 1). The seven surviving FAP patients were followed for a mean of 36 months (range 23 to 50 months) after the qualifying echocardiogram. None of these patients developed class III or class IV heart failure.

Despite the finding of identical echocardiographic appearances, a larger proportion of patients in the AL group (91%; 21 of 23) had congestive heart failure symptoms (New York Heart Association class II to IV) than in the familial group (17%; two of 12), P < 0.0001. Of these, the two FAP patients were confined to class II symptoms, whereas 14 patients in the AL group had class III-IV heart failure (table 1).

In order to investigate an inadvertent selection bias towards older or sicker AL patients we analysed survival among an additional group of AL patients who were age matched to...
the FAP group as well as to a subgroup of 16 patients without renal disease from our original study group of 24 AL patients. Twelve AL patients of mean age 47 (12) years and with a wall thickness of > 1.3 cm (mean 1.61 (0.36) cm) had a median survival of 5-5 months, and in addition only 33% of these patients survived more than one year from the date of their echocardiogram. Compared to the FAP group who had a one year survival of 92%, survival was significantly shorter in the age matched AL patients (P = 0.009). When the one year survival of the FAP group (92%) is compared to that of the 16 AL patients without renal involvement, of whom four survived one year (25%), there was a significantly worse survival in the AL patients (P = 0.0006).

LIVER TRANSPLANTATION
Liver transplants were successfully performed in six of the 12 patients with FAP. Despite the extensive myocardial infiltration on echocardiography, no patient had complications of perioperative or postoperative congestive heart failure.

Discussion
QUALITATIVE ECHO DOPPLER OBSERVATIONS
ON FAP AND AL PATIENT GROUPS
This study shows that the echocardiographic features in AL amyloidosis and FAP were present to a very similar degree in both types of amyloidosis, such that the type of amyloid could not be identified from the echocardiogram.

Klein et al have shown that heart failure and survival in AL amyloidosis is correlated with the degree of myocardial thickening seen on echocardiography.14 Our data support this finding in patients with AL amyloidosis, but show that heart failure may be minimal or absent despite extensive wall thickening in patients with the familial disease. No patient with FAP developed severe heart failure (class III or IV) and no familial patient died of chronic heart failure during the follow up period. Thus the prognosis in FAP, unlike that in light chain associated AL amyloid, cannot be predicted from the echocardiogram.

The finding of a “restrictive” Doppler pattern in three of the 12 FAP patients in the absence of heart failure appears, at first glance, difficult to explain. However a major criterion of the restrictive pattern is a small A wave in conjunction with an abbreviated E wave deceleration time.13 Patients with FAP may have other reasons for a small A wave—specifically atrial amyloid, resulting in its most extreme form in electromechanical dissociation of the atrium.22 It is possible therefore that this Doppler appearance may not reflect increased atrial afterload but rather decreased atrial contractility.

Liver transplantation, by removing the source of the mutant protein, has become an effective treatment for preventing production of familial amyloid protein (transthyretin) and arresting the disease.11 Support for the concept that wall thickening in familial amyloid is of little prognostic significance comes from the response of the six patients who underwent liver transplantation without perioperative problems of fluid balance. Our selection criteria for liver transplantation for FAP do not exclude patients who are seriously incapacitated, for example from neuropathy, inanition, or severe heart failure. It is of note that we did not have to exclude anyone in the latter category, despite the echocardiographic appearances. We do not believe that liver transplantation had any effect in prolonging survival in patients in this study, since follow up was relatively short and all patients accepted for liver transplantation were expected to live at least five years, based on clinical status.

The two groups had significantly different periods of survival. Although it might be argued that this may be related to the differences in the severity of heart failure at the time of the echocardiogram, we believe that the differences in mortality and in heart failure are intimately linked and reflect differing effects of AL and FAP amyloid infiltration of the myocardium. The onset of heart failure in patients with AL amyloid is predicted by the severity of myocardial thickening on echocardiography.14 If the same were true in FAP one would anticipate a similar prevalence of heart failure given the similar increase in left ventricular mass, yet heart failure was uncommon, never severe, and did not develop de novo during a mean follow up of 36 months in either the liver transplanted or the non-transplanted group. Furthermore, there is no described phase of myocardial infiltration in AL amyloidosis characterised by marked left ventricular thickening without significant heart failure, so it cannot be argued that the AL patients were at a later, symptomatic, stage of their cardiac disease than the familial group. We thus believe that these differences in symptoms and mortality reflect true differences in the response of the heart to deposition of amyloid fibrils in the two forms of the disease—a theory that is supported by our observations on the electrocardiographic voltage response.

ELECTROCARDIOGRAPHIC VOLTAGE AND VOLTAGE MASS
In general increased cardiac mass is associated with an increase in electrocardiographic voltage. In AL amyloidosis, there is an inverse relation between left ventricular mass and electrocardiographic voltage.23 This is due to a combination of replacement of some myocardial cells by electrically inert amyloid and destruction of many remaining myocytes. Our findings of low limb lead voltage electrocardiograms in 33% of familial and 60% of the AL patients are consistent with previous studies and represents amyloid infiltration of the myocardium.24-26 However, there were differences between the two groups which suggested that more functional myocardium remained in the FAP patients. The difference in the mean limb lead voltage between the two groups was statistically significant, as was the difference in the voltage to mass ratio. Although a low volt-
Familial and primary (AL) cardiac amyloidosis

age recording may be associated with pericardial effusions it seemed unlikely that the small effusions observed in our patients would account for the voltage differences. Indeed analysis of the electrocardiograms of those patients with \( n = 8 \) and without \( n = 16 \) effusions in the AL amyloid group showed no significant differences in voltages. Although the precise mechanism for the relatively preserved voltage in FAP is not clear, it may be postulated that it is related to different histological deposition patterns in the two disease types (for example, lesser physical distortion and destruction of the remaining myocytes in FAP), differences in the biological characteristics of the protein, or differences in myocyte response to myocardial infiltration (for example, reactive hypertrophy of the remaining myocardium in FAP). Such differences, if present, may explain the lesser degrees of heart failure and the better survival in the familial group.

**Limitations of the Study**
The number of patients with FAP and echocardiographic abnormalities studied was small and thus minor differences between groups may not have been detected. However, the very poor survival in the AL patients is consistent with similar patients in our large AL patient database and with data from other centres. The strikingly better survival in the FAP group is highly significant and consistent with other reports of a longer duration of survival in this disease compared to AL amyloidosis. As indicated in the methods section, the small number of FAP patients studied represents the total number of patients with echocardiographic abnormalities out of a series of 40 patients. Given the extreme rarity of FAP, we believe that these numbers represent one of the largest series of patients with this condition seen in the United States.

**Clinical Implications**
We have shown that despite very similar echocardiographic appearances, cardiac involvement in FAP and AL amyloidosis has a very different outcome. The clinical significance of these findings is important. If a diagnosis of amyloidosis is suspected by echocardiography and confirmed by tissue biopsy, it is critical to determine the type of amyloid, not only because the prognosis of the heart disease differs but also because the treatment of the two forms is completely different.

**Determining the type of amyloidosis**
It is rarely necessary to perform endomyocardial biopsy, and histology is usually obtained from an abdominal fat biopsy—the characteristic apple green birefringence seen when the tissue is stained with congo red and viewed under polarised light is diagnostic for systemic amyloidosis of either the AL or FAP type. After a positive tissue biopsy is obtained and if there is no family history of amyloidosis, AL amyloidosis is considered first as it is the most common type. A search for a clonal plasma cell dyscrasia is the first step. Monoclonal immunoglobulins or light chains are detected in 90% of AL patients by means of immunofixation electrophoresis of serum and urine, a more sensitive technique than simple protein electrophoresis. Patients with apparent AL amyloidosis who do not have monoclonal light chains can pose a diagnostic problem. In most of these patients, a clonal dominance of plasma cells will be identified by examining a bone marrow biopsy with immunohistochemical staining or by cellular studies employing labelled antibodies specific for human light chains. In rare cases gene rearrangement studies may be employed. When there is no evidence of a plasma cell dyscrasia, consideration should be given to another form of amyloidosis. While a family history of amyloidosis or unexplained progressive neuropathy strongly suggests FAP, a variant transthyretin is sought in all patients who do not have a plasma cell dyscrasia. Transthyretin can be identified by isoelectric focusing of the serum, which will separate variant and wild type transthyretin. The finding of a variant transthyretin in serum then prompts specific genetic testing to define the mutation precisely.

**Treatment of AL amyloidosis and FAP**
The treatment of AL amyloidosis usually involves oral chemotherapy with alkylating agents such as melphalan coupled with prednisolone. Most recently high dose intravenous melphalan has been used in an attempt to annihilate the plasma cell clone; this treatment method requires autologous stem cell rescue to repopulate the bone marrow after the chemotherapy has been given. In the case of FAP, where the abnormal protein is transthyretin, no drug treatment has proved effective and the treatment of choice is liver transplantation to remove the source of the mutant protein.

Our data indicate that in FAP there is a disparity between severe myocardial thickening and clinical features of heart failure, and they suggest that, in the absence of severe heart failure, major surgical procedures may be successfully undertaken. Thus once FAP is diagnosed the finding of echocardiographic abnormalities should not be taken as a sign of impending heart failure or as an absolute contraindication to liver transplantation. An unanswered question is the reason for the clinical differences in AL amyloidosis and FAP with severe echocardiographic abnormalities. While the electrocardiographic data suggest more residual functional myocardium in FAP, the exact reason awaits a careful historical comparison in the two patient groups and an assessment of the effects of the different biological composition of the amyloid fibrils on normal myocytes.

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