The heart in limb girdle muscular dystrophy

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A J van der Kooi, W G de Voogt, P G Barth, H F M Busch, F G I Jennens, P J H Jongen, M de Visser

Abstract

Objective—To assess the frequency, nature, and severity of cardiac abnormalities in limb girdle muscular dystrophy, and its relation to age and weakness in various genotypes.

Design—In 26 autosomal dominant, 38 autosomal recessive, and 33 sporadic strictly defined patients with limb girdle muscular dystrophy, cardiac evaluation included history, physical examination, chest x ray, electrocardiography, 24 hour ECG Holter monitoring, and echocardiography. In 35 of the 71 autosomal recessive and sporadic cases muscle biopsies were available for sarcoglycan analysis.

Main results—Dilated cardiomyopathy was present in one autosomal dominant case and in three advanced autosomal recessive or sporadic patients, of whom two were found to have a sarcoglycan deficiency. Two of these three patients and three other cases showed ECG abnormalities known to be characteristic of the dystrophinopathies. A strong association between the absence of a sarcoglycan and the presence of dilated cardiomyopathy was found (p = 0.04). In six autosomal dominant cases there were atrioventricular (AV) conduction disturbances, increasing in severity with age and in concomitant presence of muscle weakness. Pacemaker implantation was necessary in four.

Conclusions—10% of these patients had clinically relevant cardiac abnormalities. In autosomal dominant limb girdle muscular dystrophy one subtype characterised by muscle weakness and AV conduction disturbances is recognised. In the course of autosomal recessive/sporadic limb girdle muscular dystrophy, dilated cardiomyopathy may develop, probably related to deficiency of dystrophin associated proteins.

(Heart 1998;79:73–77)

Keywords: limb girdle muscular dystrophy; cardiomyopathy; AV conduction block; sarcoglycan

Cardiac involvement in muscular dystrophies is a well known complication. In Duchenne muscular dystrophy, Becker muscular dystrophy, and carriers of the dystrophin gene, assembled under the term dystrophinopathies, dilated cardiomyopathy and specific ECG abnormalities are frequent findings. In yet another type, that is, Emery–Dreifuss muscular dystrophy, conduction defects are a hallmark of the disease. There are, on the other hand, also muscular dystrophies such as Bethlem myopathy in which cardiac involvement is not present. The occasional patient with heart problems due to a dystrophic process has been described in limb girdle muscular dystrophy, a clinically and genetically heterogeneous group of disorders (table 1). However, these reports contained either anecdotal cases or small series of limb girdle muscular dystrophy patients.

The present study was part of an inventory of all known cases of limb girdle muscular dystrophy in The Netherlands, and comprises the largest group of carefully diagnosed limb patients so far. The objectives were to assess (1) the frequency, nature, and severity of cardiac abnormalities, and (2) the correlation between cardiac involvement and mode of inheritance, age, and severity of muscle weakness.

Methods

STUDY POPULATION

Ninety seven patients (59 females, 38 males) from 66 families with limb girdle muscular dystrophy were included in this cross sectional study. They participated in an explorative study of all known cases of limb girdle muscular dystrophy in The Netherlands and were subjected to a strict diagnostic procedure to exclude other possible causes of limb girdle syndrome, and to assessment of muscle strength. Dys trophy analysis of muscle tissue was performed in 42 patients from 56 families. The remaining 14 families were either multiplex or consanguineous families comprising only affected females (4), affected individuals of both sexes (2), or affected males in whom restriction fragment length polymorphisms had excluded X linked inheritance (2), and families with sporadic females (6). Where relevant, dystrophin gene screening and karyotyping was performed.

Autosomal recessive inheritance was present in 38 patients out of 23 families, autosomal dominant inheritance in 26 patients from 10 families, and 33 cases were sporadic. Age and disease duration are listed in table 2.

CARDIAC EVALUATION

Cardiac involvement was assessed by several non-invasive types of investigation. Medical and family history was taken, and a physical examination performed. Full inspiration chest radiographs were obtained in the standing or sitting position. A standard 12-lead ECG was carried out at rest in the supine position. Evaluation included a classification according to the Minnesota code.

Ambulatory ECG monitoring on a Reynolds two channel recorder for a 24 hour period of routine daily activities was performed in all
patients above the age of 10 years. A computer was used to analyse the magnetic tapes, and all tapes were read by one technician.

Cross sectional and colour flow Doppler echocardiographic investigations were performed in all patients using a Hewlett-Packard Sonos 500 and 2500 (Hewlett-Packard, Camas, Washington, USA). Echocardiographic evidence for dilated cardiomyopathy consisted of an enlarged end diastolic left ventricle (above the 95th centile, adjusted for body surface area) together with impaired systolic function. Body surface area was calculated in all patients in whom measurement of weight and length was possible. If correction by surface area was not possible, an end diastolic diameter > 60 mm was considered pathological. Systolic function was considered impaired if fractional shortening was less than 25% or if global hypokinesia on cross sectional echocardiographic examination was seen, or both. Regional wall motion abnormalities were recorded.

IMMUNOHISTOCHEMICAL STUDIES
Forty two serial cryostat sections of available muscle specimens of autosomal recessive and sporadic cases were studied for dystrophin and spectrin; 35 of these were available for α sarcoglycan screening.35

STATISTICAL ANALYSIS
χ² Tests were used to analyse the relation between severity of the cardiac disorder and age and severity of weakness, respectively. In autosomal recessive and sporadic cases the relation between the absence of α sarcoglycan and the presence of dilated cardiomyopathy was also analysed. The cut off value for the severity of weakness was the loss of independent walking, because we considered this an important indicator for the progression of the disease. Because our study was cross sectional, the statistical relations can only suggest an association.

Results
A complete cardiological work up was available in 91 out of 97 patients (94%). In three, Holter ECG was incomplete, and in three other wheelchair bound patients echocardiography was not available because of pulmonary interposition. None of the patients had symptoms related to chronic cardiac failure. All abnormal results are listed in table 2.

AUTOSOMAL RECESSIVE PATIENTS
An ambulatory 44 year old male, and a 23 year old wheelchair dependent female had an RS ratio > 1 in lead V1 on ECG. In the former, muscle sarcoglycan was normal. The latter also showed a deep lateral Q wave and in addition she had borderline echocardiographic abnormalities consisting of a slightly abnormal contraction pattern of the left ventricle. Her muscle biopsy specimen was not available for sarcoglycan analysis. One wheelchair dependent woman aged 47 had dilated cardiomyopathy with a fractional shortening of 15%. She was born from consanguineous parents, and her brother died at the age of 19 years because of cardiac failure due to dilated cardiomyopathy. Her muscle biopsy specimen was also not available for sarcoglycan analysis. An ambulatory 23 year old woman showed an abnormal contraction pattern of the interventricular septum. Muscle tissue was available from one of her siblings for sarcoglycan analysis, and she had a reduction of α sarcoglycan. In a 46 year old wheelchair dependent patient with α sarcoglycan deficiency, no cardiac abnormalities were found (see also table 3).

SPORADIC PATIENTS
An ambulatory 44 year old male with normal sarcoglycan and a 17 year old wheelchair dependent girl showed an RS ratio > 1 in lead V1 on ECG. The latter, and a 27 year old female who had great difficulty walking, also had a narrow Q in the lateral leads, and they both had dilated cardiomyopathy. Both these patients turned out to have α sarcoglycan deficiency, but normal dystrophin. Another female aged 20 was found to have an abnormally contracting interventricular septum; her muscle biopsy was not available for sarcoglycan analysis. Of three other wheelchair dependent patients with α sarcoglycan deficiency, one

### Table 1 Classification of limb girdle muscular dystrophy

<table>
<thead>
<tr>
<th>Subtype (chromosomal localisation)</th>
<th>Gene product</th>
</tr>
</thead>
<tbody>
<tr>
<td>Autosomal dominant, LGMD1 (5q)</td>
<td>Not identified</td>
</tr>
<tr>
<td>Autosomal recessive, LGMD2 (1q)</td>
<td>Not identified</td>
</tr>
<tr>
<td>LGMD2A (15q)</td>
<td>Calpain (57)</td>
</tr>
<tr>
<td>LGMD2B (2p10)</td>
<td>Not identified</td>
</tr>
<tr>
<td>LGMD2C (13q14)</td>
<td>γ Sarcoglycan (52)</td>
</tr>
<tr>
<td>LGMD2D (17q)</td>
<td>α Sarcoglycan (52)</td>
</tr>
<tr>
<td>LGMD2E (4q)</td>
<td>β Sarcoglycan (52,54)</td>
</tr>
<tr>
<td>LGMD2F (5q)</td>
<td>δ Sarcoglycan (52)</td>
</tr>
</tbody>
</table>

*One patient can show more than one ECG or echocardiographic feature; † atrial fibrillation, extrasystoles, and atrial rhythm. AV, atroventricular.

### Table 2 Results of cardiological investigations

<table>
<thead>
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rhythm, and a ventricular extrasystole were noted (paper speed 25 mm/s).

atrial fibrillation to sinus rhythm, total atrioventricular block with junctional escape
muscle weakness and atrioventricular conduction disturbances. After electroconversion from

Figure 1 Rhythm strip of a member of the autosomal dominant families characterised by
The heart in limb girdle muscular dystrophy

of dilated cardiomyopathy (table 3; p = 0.04).

had no cardiac abnormalities.

STATISTICAL ANALYSIS

χ² Tests showed a strong association between the absence of α sarcoglycan and the presence of
dilated cardiomyopathy (table 3; p = 0.04).

AUTOSOMAL DOMINANT PATIENTS

In six of 15 cases from two families with auto-
somal dominantly inherited limb girdle muscu-
lar dystrophy, atrioventricular conduction dis-
turbances were seen (fig 1), in the concomitant
presence of muscle weakness. Because none of
the patients had lost the ability to walk
independently, a relation with weakness could
not be established. χ² Testing showed a signif-
ificant relation between the severity of atrioven-
tricular conduction disturbances and age
(p < 0.001). These patients have been
described in detail elsewhere.13 Two had first
degree, two second degree, and two total atrio-
ventricular block. Two patients had a history of
syncopal attacks and dizziness, requiring im-
plantation of a pacemaker at the age of 62 and
40 years, respectively. During our study two
other patients developed symptomatic atrio-
ventricular block. (Paroxysmal) atrial fibrillation
was found in four autosomal dominant pa-
tients. One was found to have dilated cardio-
myopathy, with an end diastolic diameter of
70 mm and poor contraction of all walls. In
addition, two other patients showed abnormal
contractions of the interventricular septum.
Atrioventricular block was not associated with
echocardiographic abnormalities.

In three cases from other autosomal domi-
nant families, left ventricular hypertrophy was
encountered. These patients had hypothy-
roidism, hypertension, and aortic valve stenosis
with a peak systolic gradient of 40 mm Hg,
respectively.

Discussion

There are five recent small reports on cardio-
logical involvement in limb girdle muscular
dystrophy. In only three of these4–7,30 strict diag-
nostic criteria for limb girdle muscular dystro-
phy have been used. In the study by Stübbgen,34
which comprised 20 limb girdle muscular dys-
trophy patients, none of the patients had symp-
tomatic heart disease. However, subclinical
non-specific cardiac abnormalities were de-
tected in 80%. Whether these could be
ascribed to a dystrophic process of the heart
remains unclear. In the second study,35 describ-
ing 12 patients, two brothers with α sarcogly-
can (previously called adhalin) deficiency and
dilated cardiomyopathy have been mentioned.
The third report describes a patient with
dilated cardiomyopathy and α sarcoglycan
deficiency who had previously been given a
diagnosis of Becker muscular dystrophy.36 The
remaining two reports described a patient with
an impaired left ventricle function without
associated dilatation,37 and three adult sisters
with dilated cardiomyopathy.38 Recent detailed
clinical descriptions of the different subtypes
of limb girdle muscular dystrophy mention the
absence of cardiac abnormalities in type 2B,38
the very rare presence of abnormalities in type
2A,39 40 and a high frequency of cardiac anoma-
lies in type 2C.41 These reports suggest that
deficiency of one of the components of the
 dystrophin associated glycoprotein complex
might cause dilated cardiomyopathy. The fact
that the sarcoglycan complex is expressed in
the heart42 43 supports this view.

In this study, the largest so far conducted in
which all patients have met strictly defined cri-
teria for a diagnosis of limb girdle muscular
dystrophy, various cardiac abnormalities were
encountered. In our patient population dilated
cardiomyopathy was present in three of 71
autosomal recessive or sporadic patients, com-
parable with the prevalence in dystrophinopa-
thies. They were more advanced, but otherwise
healthy, cases. Two were sporadic female
patients, aged 17 and 27, respectively, both
with α sarcoglycan deficiency. Although the
numbers were small, χ² testing nevertheless
showed a strong association between the
absence of α sarcoglycan and the presence of
dilated cardiomyopathy. Direct mutation
analysis of the genes encoding the four
sarcoglycan subunits is needed to pinpoint the
precise genotype, since absence of α sarcogly-
can is always observed, no matter which of the
sarcoglycans is lacking.30 These studies are
currently in progress and may give more insight
into the mechanism responsible for the involve-
ment of cardiac muscle in some patients with
limb girdle muscular dystrophy.

Interestingly, dystrophinopathy specific ECG
abnormalities, including increased R waves in
lead V1, an RS ratio > 1 in V1, and abnormal Q
waves in the lateral leads,4 5 were found in
the two sporadic females with dilated cardiomy-
opathy, and in three other patients (including
two males). One of these three showed border-
line echocardiographic abnormalities, consisting
of a slightly abnormal contraction pattern of
the left ventricle. Conceivably these ECG
abnormalities may be considered an early stage
in the development of dilated cardiomyopathy in
limb girdle muscular dystrophy.

Table 3 Relation of sarcoglycan deficiency to dilated cardiomyopathy in autosomal recessive and sporadic patients

<table>
<thead>
<tr>
<th>Sarcoglycan</th>
<th>Present</th>
<th>Absent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Absent</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>Normal</td>
<td>0</td>
<td>28</td>
</tr>
<tr>
<td>Unknown</td>
<td>1</td>
<td>20</td>
</tr>
</tbody>
</table>

Figure 1 Rhythm strip of a member of the autosomal dominant families characterised by
muscle weakness and atrioventricular conduction disturbances. After electroconversion from
atrial fibrillation to sinus rhythm, total atrioventricular block with functional escape
rhythm, and a ventricular extrasystole were noted (paper speed 25 mm/s).
In families with autosomal dominantly inherited limb girdle muscular dystrophy, age related atrioventricular conduction disturbances were seen. Several patients developed symptomatic atrioventricular block. We strongly advocate 24 hour ECG recording, when ECG abnormalities suggesting atrioventricular conduction disturbances are present, in order to identify atrioventricular block at an early stage. Regular follow up examinations and implantation of a pacemaker if the patient becomes symptomatic are preventive measures. To a certain extent these families resemble Emery-Dreifuss muscular dystrophy, but the absence of early contractures and rigid spine, the limb girdle distribution of weakness, and the mode of inheritance clearly distinguish these disorders. The occurrence of Wolff-Parkinson-White syndrome in one patient is probably of no significance because of its relatively high prevalence of 0.15% in the normal population. Hypertrophy of the left ventricle was found in 10% of our patients we found hypertrophy of the left ventricle with cardiac valve stenosis. Whether this finding is indicative of involvement of the heart as part of the dystrophic process is as yet uncertain.

CONCLUSIONS

In summary, in 10% of our patients we found clinical evidence of cardiac involvement and that is, atrioventricular conduction disturbances related to age in an autosomal dominant subtype and dilated cardiomyopathy in some advanced cases of the autosomal recessive/sporadic group. This refutes the previous notion that cardiological abnormalities in limb girdle muscular dystrophy are uncommon.

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