Case reports


Kearns’ syndrome, a new form of cardiomyopathy

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A 19-year-old boy was admitted with Adams-Stokes syndrome. Subsequent examination revealed bilateral exophthalmoplegia, ptosis, and retinal pigmentary change. A pacemaker was implanted because of complete heart block. Only 12 similar cases have been previously described.

The detection by Meerschwan of abnormalities in skeletal muscle in patients with hypertrophic obstructive cardiomyopathy has raised the question that this disease may represent a more generalized muscle disorder, but the matter is still unresolved (Goodwin, 1970; Meerschwan, 1969).

Cardiomyopathy is a recognized entity in muscle dystrophies and myotonia dystrophica, where there is generalized degeneration of skeletal muscles (Welsh, Lynn, and Haase, 1963).

Sandifer described cardiomyopathy in association with external ophthalmoplegia (1946). Later, Kearns and Sayre (1958), and Kearns (1965) described 9 cases with external ophthalmoplegia, retinal pigmen-
tary change, and complete heart block. The purpose of this communication is briefly to review the salient features of this interesting syndrome described by Kearns. Furthermore, haemodynamic data not hitherto reported, in a patient thought to have this syndrome, are presented. Kearns’ syndrome deserves attention in cardiology because of the presence of complete heart block.

Case report

In 1963 a 12-year-old boy was referred to the Eye Hospital of the University of Utrecht because of embarrassing ptosis, which was surgically corrected. Six years later he was referred to a neurologist because of two ‘black-outs’ on the road. An electroencephalogram showed no evidence of epilepsy or other abnormalities.

In March 1970, when 19 years old, he was admitted to the Cardiology Unit of the University Hospital in Utrecht, with a history of transient loss of consciousness while walking. Physical examination revealed a normally-developed boy with divergent strabismus, generalized limitation of all eyeball movements, and conspicuous ptosis; dark adaptation was normal. Fundus oculi showed peripapillary pigment deposit giving a halo appearance but no evidence of retinitis pigmentosa.

In the cardiovascular system the heart rate was 44-72/min, and the blood pressure 130/60 mmHg. The first sound was normal and followed by a grade 1/4 proto-
systolic murmur without a definite point of maximal intensity. There was wide splitting of the second heart sound, both components being of equal intensity. At the apex, a third heart sound was heard. The rest of the physical examination revealed no abnormalities. There was no evidence of myasthenia, myotonia, or muscular hypertrophy elsewhere.

Laboratory findings Blood group A rhesus positive. The blood picture and ESR were normal. Twenty-four-hour creatinine excretion in urine 1.2 mg (low). Blood urea 52 mg/1000 ml, creatinine 11 mg/1000 ml, serum electrolytes, protein-bound iodine, liver function tests, lipid spectrum, blood sugar, serum aspartate amino-
transferase and alanine transferase, lactase dehydrogen-
ase, and creatine phosphokinase enzymes were all normal. Anti-heart muscle antibody was negative, anti-
skeletal muscle antibody doubtfully positive. Toxo-
plasma dye test positive in 1/512, and complement-
fixation test positive in 1/8. Rectal biopsy showed no evidence of amyloidosis.

Electrocardiogram (Fig. 1) This showed sinus rhythm at 60 a minute: PR interval 0-20 sec, QRS duration 0-12 sec. There were periods of first-degree and second-degree heart block as well as periods of sinus arrest. The QRS complexes showed right bundle-branch block configuration with left axis deviation, compatible with partial bilateral bundle-branch block. The possibility of biventricular hypertrophy could not be excluded. The chest x-ray revealed slight left ventricular enlarge-
ment.

Jugular venous tracing Y-descent deeper than x-descent.

Carotid pulse tracing Normal. On cardiac catheter-
izatization – all intracardiac and intravascular pressures
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were normal. The oxygen saturations were normal. The left ventricular end-diastolic pressure was 8 mmHg and peak dp/dt value 980 mmHg/sec. The cardiac index was 3.6 l/min. There was no evidence of Ebstein's disease or intracardiac shunt.

**Cineangiocardiography** Injection of dye into the left ventricle showed slight mitral reflux during ventricular diastole. There were no premature beats during dye injection and no signs of hypertrophic obstructive cardiomyopathy or evident left ventricular wall thickening. On contrast injection into the root of the aorta no coronary arterial obstruction could be seen.

**On audiometry** a bilateral perceptive dip was demonstrable and there was no labyrinthine response to chloretic.

**Cerebrospinal fluid** Protein 124 mg/100 ml, on electrophoresis gammaglobulin 15 per cent, leucocytes 96, later 3. Erythrocytes nil.

**Electroencephalogram** Irregular with diffuse changes (abnormal).

**Electromyogram** (external ocular rectus muscle) Myopathic pattern.

**Ocular fluorescein angiogram** Besides tapetoretinal pigmentary changes there was also a tapetochoroidal component present especially in the peripapillary area. These findings do not fit into the classical tapeto-choroidal dystrophies (Fig. 2).

**FIG. 1** Electrocardiogram. Record A shows first-degree heart block and period of sinus arrest. There is an extreme left axis deviation. Record B shows a complete right bundle-branch block and a second-degree heart block.

**FIG. 2** Fluorescein ocular angiogram: showing peripapillary tapetochoroidal pigmentary change.
Progress
During cardiac catheterization the patient developed Adams-Stokes attacks due to ventricular asystole and was temporarily paced with a transvenous bipolar catheter. The heart block did not respond to atropine or isoprenaline therapy. A pacemaker (Cordis) was later implanted. His ophthalmoplegia and ptosis have remained stationary since 1963. He is doing well up to the time of publication of this paper.

Discussion
Summarizing, this patient has partial bilateral bundle-branch block with Adams-Stokes syndrome, an abnormal jugular venous tracing, slight diastolic mitral reflux, bilateral external ophthalmoplegia, bilateral ptosis, peripapillary retinal pigmented change, asymptomatic labyrinthine disturbance, audiometric perceptual dip, raised gammaglobulin in the cerebrospinal fluid, an abnormal electroencephalogram, a myopathic-pattern electromyogram, and atypical tapetochoroidal dystrophy on fluorescein angiography. Chronic progressive external ophthalmoplegia was for the first time called oculomypathy in a classic paper by Kiloh and Nevin (1951), stressing the myopathic nature of the disease. In their series there was no cardiomyopathy. The first authentic association of external ophthalmoplegia and cardiomyopathy was described by Sandifer (1946). Marinesco (1910), in his monograph on diseases of muscle, asserts that of the muscles which offer extraordinary resistance to involvement by muscular dystrophy, the heart is the most notable. Kearns and Sayre (1958) published a series of nine cases under the heading: 'External ophthalmoplegia, pigmentary degeneration of the retina and cardiomyopathy: a new recognized syndrome'. Five out of these nine cases had all the foresaid three components; frequent associations were peripheral and facial muscle weakness, deafness, small stature, abnormal electroencephalogram, and increased protein content of the cerebrospinal fluid. The age of onset varied from 3 to 25 years, sex incidence being equal. He concluded that external ophthalmoplegia, cardiomyopathy, and peripheral muscle weakness were all a result of a generalized muscle disorder. Abnormal electroencephalogram and increased protein in the cerebrospinal fluid suggest central nervous involvement. He asserted that the association between ophthalmoplegia, retinal pigmentary degeneration, and cardiomyopathy does not appear to be merely a chance occurrence but a syndrome.

The patient described here has enough clinical features to justify a diagnosis of Kearns' syndrome. It is interesting to note that the cardiac output and end-diastolic pressure in the left ventricle are normal. Furthermore, in the jugular venous tracing the y-descent is deeper than the x-descent, which is corroborated in the right atrial pressure curve. Normally the x-descent is deeper than the y-descent in jugular venous tracing. A y-descent deeper than the x-descent has been described by Hartman (1960) in tricuspid insufficiency, atrial fibrillation, operated cases of atrial septal defect, and ventricular septal defect.

In five cases reported by Kearns, four had right bundle-branch block with left axis deviation. In the present case also a similar combination was found, compatible with a diagnosis of bilateral partial bundle-branch block (Rosenbaum, 1970). In this patient mitral reflux started in mid-diastole and continued up to the beginning of the next ventricular systole. This is best explained as described by Rutishauser et al. (1966) who reported atrogenic reflux in patients with complete atrioventricular block. This was explained on the basis of ventricular to atrial pressure gradient created by atrial relaxation.

This case is apparently the first of this syndrome in which the peripapillary retinal pigmentary change was analysed with the help of fluorescein angiography. There was striking choroidal fluorescence present which formed the background for fine piles of pigment. Retinal vascular filling and emptying were normal. This peripapillary tapetochoroidal pigmentary change is atypical for any of the tapetochoroidal dystrophies so far described.

Kearns forecast an ominous prognosis in these cases due to the problems of complete heart block. Now with modern pacing techniques one can offer new hope to these patients. The present case is doing well two years after implantation of a cardiac pacemaker.

Electrocardiographic features such as right bundle-branch block with left axis deviation and first-degree heart block in a young subject with ptosis should arouse the suspicion of a generalized syndrome.

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References


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