Case Report

A patient with cytoplasmic body myopathy presented muscle hypotonia from birth and developed progressive muscular atrophy and weakness, scoliosis, contracture of joints and cardiorespiratory failure. At the age of 17, he died of heart failure. Post mortem examination revealed severe hypertrophy of cardiac walls and generalized muscular atrophy. Microscopic examination showed many cytoplasmic bodies in skeletal muscle fibers and myofiber disarray in myocardium. No cases of cytoplasmic body myopathy with hypertrophic cardiomyopathy have been reported previously. It is suggested that the Z-line component is related to the formation of the cytoplasmic body in skeletal muscle and disarray in the cardiac muscle.

Introduction

Cytoplasmic body myopathy is one of the very rare congenital myopathies which is characterized by many cytoplasmic bodies in skeletal muscle fibers (1-14). Clinically, cytoplasmic body myopathy reveals rather heterogeneous pictures as to heredity, age of onset, clinical symptoms and prognosis (8, 11). Only seven cytoplasmic body myopathy patients with cardiac disorders have been reported in the literature (2, 5, 7-9), and no cytoplasmic body myopathy patient with hypertrophic cardiomyopathy (HCM) has ever been documented. Here, we report a cytoplasmic body myopathy patient with severe muscular atrophy, scoliosis, joint contracture and HCM, and discuss the relation between cytoplasmic bodies in skeletal muscles and myofiber disarray in the cardiac muscle.

Case Report

The patient is a 17-year-old Japanese boy. An aunt of his father suddenly died of unknown cause at the age of 60 and a cousin of his father had idiopathic scoliosis, but there was no family history of neuromuscular disorders nor consanguinity. He was the product of a full-term pregnancy with normal delivery. However, he presented mild muscular hypotonia since birth and, subsequently, delayed motor developmental milestones with progressive muscle wasting and weakness were noted. Scoliosis and contracture of joints appeared when he was 4 years old. He suffered from dyspnea on exercise and palpitation from the age of 12 and was admitted to our hospital at the age of 17.

On admission, his body weight was 33 kg and his height was 138 cm. The pulse was irregular at 100/min and his blood pressure was 110/70 mmHg. He was an intelligent boy. Cyanosis, jugular vein enlargement and generalized edema were found. Severe scoliosis, deformity of thorax, and contracture of limb joints were seen, and generalized muscular atrophy and weakness were observed (Fig. 1). Deep tendon reflexes were decreased in all limbs, and sensory disturbance or any symptoms suggestive of central nervous system involvement were lacking.

Routine blood studies showed the following abnormalities: GOT 61 IU/L (normal, 12-37), LDH 293 IU/L (114-220), γ-GTP 131 IU/L (8-50), CK 469 IU/L (43-272), aldolase 7.7 IU/L (0.5-3.1). Arterial blood gas analysis at resting state showed severe hypoxia and hypercapnia (pH 7.374, PCO₂ 71.7 mmHg, PO₂ 30.9 mmHg, SaO₂ 57.4%). Cardiomegaly (cardiothoracic ratio 82.4%) with pulmonary congestion was seen on a chest roentgenogram. Pulmonary function tests revealed restrictive ventilatory insufficiency [forced vital capacity (FVC) 500 ml, %FVC 15.8%]. Electromyography showed polyphasic potentials...
CBM with HCM

Fig. 1. General appearance at the time of admission (17 years old). Severe muscle atrophy, deformity of thorax and scoliosis are observed.

of small amplitude but no myogenic pattern.

Electrocardiogram (ECG) revealed sinus tachycardia, left axis deviation, first degree atrioventricular block, incomplete right bundle branch block, right atrial enlargement and right ventricular hypertrophy. Holter ECG showed frequent premature ventricular contraction, premature supraventricular contraction, and paroxysmal supraventricular tachycardia. Echocardiography disclosed slight pericardial effusion, marked enlargement of both atriums, and hypertrophy of both ventricular walls. Doppler echocardiography showed a slight tricuspid regurgitation. Cardiac catheterization revealed severe pulmonary hypertension and both ventricular filling pressures were elevated (Table 1). Three months later, he died of heart failure.

Table 1. Cardiac Catheterization Data of This Patient

<table>
<thead>
<tr>
<th>Catheterization data</th>
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<tbody>
<tr>
<td>RA</td>
<td>24 mmHg</td>
</tr>
<tr>
<td>RV</td>
<td>63/EDP 28 mmHg</td>
</tr>
<tr>
<td>PA</td>
<td>74/48 (54) mmHg</td>
</tr>
<tr>
<td>PCW</td>
<td>27 mmHg</td>
</tr>
<tr>
<td>CO</td>
<td>2.51 L/min</td>
</tr>
<tr>
<td>CI</td>
<td>2.49 L/min/m²</td>
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RA: mean right atrial pressure, RV: right ventricular pressure, PA: pulmonary artery pressure, PCW: mean pulmonary capillary wedge pressure, CO: cardiac output, CI: cardiac index.

Autopsy findings

At autopsy, there were no remarkable pathologic changes except for the skeletal muscles and heart. Frozen sections of 8 μm thickness were prepared from diaphragm, biceps brachii, quadriceps femoris, iliopsoas and paravertebral muscles. All muscle specimens showed mild variation in fiber size, increase of internal nuclei and many vacuoles. The most notable feature was the presence of intracytoplasmic inclusions in some degenerated muscle fibers: these cytoplasmic bodies were round or oval, and appeared reddish with HE stain (Fig. 2A) and purplish with modified Gomori-Trichrome stain (Fig. 2B), but did not show any histochemical reactivities for either NADH-tetrazolium reductase or routine ATPase at pH 9.4. Dystrophin immunostaining was performed using 1:500 mouse monoclonal antibody 4C5 raised against a C-terminal peptide of dystrophin (15), revealing the normal localization of this protein in the muscle fibers examined. Formalin-fixed muscle tissues were post-fixed with 1% osmium tetroxide, embedded in epoxy resin, and used for conventional electron microscopic observations. The core of the cytoplasmic bodies showed a fine filamentous appearance intermixed with dense amorphous material and the electron density of these bodies closely resembled that of Z-line (Fig. 3A). Moreover, some cytoplasmic bodies contained structures similar to deformed Z-lines (Fig. 3B) and the periphery of

![Figure 2A](image1.png)
![Figure 2B](image2.png)

Fig. 2. Representative histological findings of muscles. A) Diaphragm, HE stain, x150. B) Right biceps brachii, Modified Gomori-trichrome stain, x100. Many cytoplasmic bodies are visible in muscle fibers. Arrows indicate typical cytoplasmic bodies in muscle fibers.
Cytoplasmic bodies are unique sarcoplasmic inclusions of the skeletal muscle fibers (16), which can be found in various neuromuscular disorders, including inflammatory myopathy, myotonic dystrophy, progressive muscular dystrophy, periodic paralysis and mitochondrial myopathies (17). The term cytoplasmic body myopathy has been applied when many muscle fibers contain cytoplasmic bodies but no other definitive diag-

Discussion

Cytoplasmic bodies are unique sarcoplasmic inclusions of the skeletal muscle fibers (16), which can be found in various neuromuscular disorders, including inflammatory myopathy, myotonic dystrophy, progressive muscular dystrophy, periodic paralysis and mitochondrial myopathies (17). The term cytoplasmic body myopathy has been applied when many muscle fibers contain cytoplasmic bodies but no other definitive diag-
nosis could be made (8). Twenty-five patients of cytoplasmic body myopathy have been reported in the literature (1–14), but the clinical pictures of cytoplasmic body myopathy vary as to heredity, onset, clinical symptoms and prognosis (8, 11). This disorder, therefore, seems to encompass several different etiologies.

Chou and Mizuno reported that experimentally, spheroid cytoplasmic bodies are induced by local tetanus in rat plantaris muscle and these spheroid cytoplasmic bodies stain positively for desmin and actin immunohistochemically (18). Other studies revealed that cytoplasmic bodies in cytoplasmic body myopathy also stain positively for desmin (19), actin (14) and α-actinin, which are all contractile proteins known to be linked at the level of the Z-line. On electron microscopy, MacDonald and Engel reported that cytoplasmic bodies appear to originate from the Z-line, because they are located in the same plane as the Z-line, and their dense cores appear continuous with the Z-line (17).

Cytoplasmic body myopathy with cardiac dysfunctions has been reported in only seven patients (2, 5, 7–9) (Table 2). Almost all the patients of cytoplasmic body myopathy with cardiopathy presented a progressive muscular atrophy and poor prognosis, as in the present patient, but the details of the cardiac disorders were not described. Clinical symptoms of heart failure and abnormalities of ECG or echocardiography in these patients do not necessarily indicate primary disturbance of cardiac muscle, since severe weakness of respiratory muscles and/or marked deformity of the thorax also cause congestive heart failure. In the present patient, marked hypertrophy of both ventricular walls on echocardiography, diastolic dysfunction in cardiac catheterization, and myofiber disarray of myocardium on light microscopic examination support primary disturbance of cardiac muscle, that is, HCM.

It is intriguing in this patient that cytoplasmic bodies in skeletal muscles and HCM in cardiac muscle coexisted. These findings suggest that an abnormality of some muscle components which is expressed both in skeletal muscle and cardiac muscle might be the basic defect in this patient. The molecular genetic background of cytoplasmic body myopathy remained unidentified while data on the molecular genetics of familial hypertrophic cardiomyopathy (FHC) are accumulating. FHC, which is transmitted as an autosomal dominant trait, is a major cause of sudden death in the young. The locus of genetic abnormality in this disease was first mapped to chromosome 14q11–q12 (20) and revealed that the faulty gene codes for β myosin heavy chain, one of the major contractile proteins of the human cardiac muscle and slow skeletal muscle fibers (21). Subsequent studies revealed that mutations in the α-tropomyosin gene and cardiac troponin T gene cause FHC linked to chromosome 15q2 and chromosome 1q3 (22–24). Other loci were mapped to chromosome 11 (25), 18q (26), 16 (27) and 2q (28). It is not known what these genes code for. However, it is suggested that they encode other contractile proteins, which are expressed mainly in the cardiac muscle (α myosin heavy chain, cardiac myosin light chain 1 and 2, cardiac α actin, cardiac troponin C, I and others). Each of these protein isoforms has a role in the structure or contractile function of cardiac muscle.

We do not know the molecular basis of HCM in this case, although there is a possibility that an abnormality of a muscle-specific gene caused the cytoplasmic bodies in the skeletal muscle and the disarray of cardiac muscle fibers. A defect in a gene-coding for α-Z-line component could cause an abnormality in the muscle cell structure. Indeed, over-expression of truncated α-actinin formed a hypertrophied Z-line and nemaline-like bodies in cultured myotubes (29). In the present patient, it

<table>
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<tr>
<th>Table 2. Summarized Clinical Pictures of Patients with Cytoplasmic Body Myopathy Showing Cardiac Disorders</th>
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<tr>
<td><strong>Kinoshita (2) et al</strong></td>
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<tr>
<td><strong>Sex</strong></td>
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<tr>
<td><strong>Age at onset</strong></td>
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<tr>
<td><strong>Mode of inheritance</strong></td>
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<tr>
<td><strong>Muscle atrophy</strong></td>
</tr>
<tr>
<td><strong>Respiratory failure</strong></td>
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<td><strong>Contraction of joints</strong></td>
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<td><strong>Scoliosis</strong></td>
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<td><strong>Abnormality of</strong></td>
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<td><strong>ECG</strong></td>
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<tr>
<td><strong>UCG</strong></td>
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<tr>
<td><strong>catheterization</strong></td>
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<tr>
<td><strong>histology of myocardium</strong></td>
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<td><strong>Serum CK</strong></td>
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<td><strong>Prognosis</strong></td>
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<td><strong>Autopsy</strong></td>
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AD: autosomal dominant trait, AR: autosomal recessive trait, /: not recorded, ECG: electrocardiogram, UCG: ultrasonic cardiography.
was not clearly demonstrated that the cytoplasmic bodies originated from Z-line. However, cytoplasmic bodies and HCM can be related to the Z-line structure and some Z-line components, such as α-actinin (30) and desmin which are expressed both in skeletal and cardiac muscles. Thus, the abnormality of the Z-line component might be responsible for Z-line-related cytoplasmic bodies and HCM.

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References

19) Osborne M, Goebel HH. The cytoplasmic bodies in a congenital myopathy can be stained with antibodies to desmin, the muscle-specific intermediate filament protein. Acta Neuropathol (Berl) 62: 149, 1983.