THE FIRST FAMILY WITH KENNEDY DISEASE REPORTED FROM IRAN

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Kennedy disease is a rare X-linked, recessively-inherited disorder, which is diagnosed on the basis of clinical and electromyographic (EMG) findings, and more recently by polymerase chain reaction (PCR). The disease has been reported from around the world.

Here, we present 8 members of an Iranian family afflicted with the disease. The diagnosis was supported by PCR findings in 3 while in 5 cases the diagnosis was made based on clinical and EMG findings only.

The patients, all males, ranged in age from 20 to 85 years with the onset of gynecomastia during early adulthood and other more specific symptoms during the 5th and 6th decades of life with gradual onset and progression. The symptoms and signs included cramps, generalized weakness, dysphagia, dysphonia, tongue atrophy, tremor, generalized muscle atrophy, fasciculation, areflexia, and impotence. Four of them had hyperlipidemia and 3 had increased levels of blood glucose including one with frank diabetes mellitus. PCR, performed in 3 of the patients, showed an increase in cytosine-adenine-guanine (CAG) repeats on the long arm of X chromosome. Two heterozygote female carriers were diagnosed by the same test.

It is important to consider Kennedy disease in the differential diagnosis of lower motor neuron disorders considering its rather benign course and the need for genetic consultation for carriers.

Keywords: Bulbospinal muscular atrophy • Kennedy disease • Kennedy syndrome

Introduction

Kennedy disease is a distinct, X-linked recessive disorder characterized by progressive wasting in the proximal muscles of the lower extremities and bulbar palsy. There is often a longstanding history of muscle cramps preceding the onset of weakness in the fifth decade of life. The presence of gynecomastia, postural tremor of upper limbs, perioral fasciculation, invariable involvement of tongue, and sensory impairment in the lower limbs are the clues to diagnosis.1,2

A mutation in the coding region of the androgen-receptor gene results in a variable increase in the number of cytosine-adenine-guanine (CAG) trinucleotide tandem repeats at this site. This can be detected by polymerase chain reaction (PCR) performed on DNA from peripheral leukocytes.3,4

Kennedy et al first reported this disease in 1968. Since then, there have been a handful of reports from around the world. To the best of our knowledge, this is the first report of Kennedy disease from Iran.5–9

It is important to differentiate Kennedy disease from a more malignant, more frequent, and very similar disease, namely amyotrophic lateral sclerosis (ALS), because of the psychological implications of the diagnosis of the latter for the patient, his/her family, and the physician.
Case Report

A family (Figure 1) with 40 members in generations came to our notice when one member (the proband, case III-11) presented with the classic symptoms of the disease. He was a 58-year-old man with a history of gynecomastia for about 20 years and impotence for 8 years. He began to experience weakness, fasciculations, and cramps from the age of 43 years with a very slow progression rate. A few years later, he developed mild bulbar palsy and limb tremor. His neurologic examination showed prominent perioral and tongue fasciculations with atrophic changes. He had dysphonia, mild proximal weakness with atrophy, diffuse fasciculations and areflexia in all limbs, waddling gait, fine postural tremor in hands, and no sensory signs. The systemic examination was unremarkable except for the presence of gynecomastia. He had hyperglycemia, mild hyperlipidemia, and an increase in the serum level of creatine phosphokinase. His electromyographic exam showed prominent neurogenic changes in all the examined muscles including bulbar muscles (Figure 2). The nerve conduction studies were unremarkable except for slightly decreased sensory nerve action potential amplitudes. PCR was performed and an increased number of CAG repeats were observed as an abnormal band compared to a normal subject (Figure 3).

Seven other members of this family were diagnosed as Kennedy disease based on clinical and electromyographic examinations (in 2 of whom PCR supported the diagnosis). Findings are

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**Figure 1.** Four-generation pedigree of the index case.  
- Healthy male  
- Healthy female  
- Female  
- Male with the disease  
- Carrier (proband).

**Figure 2.** PCR in 2 carriers [IV-2 and IV-13 (right)], 2 patients [III-1 and III-8 (left)] normal person, and marker (middle).

**Figure 3.** Electromyographic tracing from the index case showing prominent neurogenic changes.  
[Contraction (right deltoid muscle), sensitivity = 1 mv]
The first family with Kennedy disease reported from Iran presented in Table 1. A certain degree of genetic anticipation was observed. The disease had a variable age of onset and tempo of progression within a single generation. Hyperlipidemia was more prominent than hyperglycemia. PCR was performed on 2 of the suspected carriers (cases V-2 and IV-13) and was positive in both.

<table>
<thead>
<tr>
<th>Case</th>
<th>Age (yr)</th>
<th>Age of onset</th>
<th>Symptoms and signs</th>
<th>Laboratory findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>III-11 (proband) 58</td>
<td>4th decade</td>
<td>Gynecomastia, cramps, generalized weakness, dysphagia, tongue atrophy, tremor, atrophy, fasciculation, areflexia, and impotence</td>
<td>CPK ↑, LDH ↔, cholesterol ↑, triglyceride↑, neurogenic changes on EMG, and CAG repeats on Xq in PCR ↑</td>
<td></td>
</tr>
<tr>
<td>II-2 85</td>
<td>6th decade *</td>
<td>Gynecomastia, atrophy, dysphagia, and gynecomastia</td>
<td>Cholesterol ↑, triglyceride ↑, and blood glucose ↑</td>
<td></td>
</tr>
<tr>
<td>II-6 77</td>
<td>6th decade *</td>
<td>Gynecomastia, atrophy, dysphagia, and gynecomastia</td>
<td>Blood glucose ↑</td>
<td></td>
</tr>
<tr>
<td>III-1 61</td>
<td>6th decade *</td>
<td>Gynecomastia, weakness, atrophy, dysphagia, and tremor</td>
<td>CPK ↑, LDH ↔, cholesterol ↑, triglyceride↑, blood glucose ↔, and CAG repeats on Xq in PCR ↑</td>
<td></td>
</tr>
<tr>
<td>III-4 46</td>
<td>——</td>
<td>Milder, similar to III-1</td>
<td>——</td>
<td></td>
</tr>
<tr>
<td>III-8 67</td>
<td>5th decade</td>
<td>Gynecomastia, DM, back pain, generalized weakness, dysphagia, dysphonia, tremor, atrophy, areflexia, and impotence</td>
<td>CPK ↑, LDH ↔, cholesterol ↑, triglyceride↑, and CAG repeats on Xq in PCR ↑</td>
<td></td>
</tr>
<tr>
<td>III-10 63</td>
<td>4th decade</td>
<td>Gynecomastia, generalized weakness, dysphonia, and fasciculation</td>
<td>——</td>
<td></td>
</tr>
<tr>
<td>IV-17 20</td>
<td>——</td>
<td>Gynecomastia (only)</td>
<td>——</td>
<td></td>
</tr>
</tbody>
</table>

CPK = creatine phosphokinase; LDH = lactate dehydrogenase; EMG = electromyography; PCR = polymerase chain reaction; DM = diabetes mellitus; ↑ = increased; ↔ = normal; * The onset of gynecomastia was earlier.
Discussion

Like most other reported cases of the disease, our cases showed an X-linked recessive pattern of inheritance within the family. The clinical pattern of presentation and the laboratory findings (except for the prominence of hyperlipidemia) were also compatible with other reported cases. These findings show that this rather benign disease (compared to the more malignant and related disease, ALS) whose diagnosis is important from the standpoint of prognosis and genetic consultation can be easily diagnosed (at least clinically) considering the classical presenting features.

References