Isaacs’ Syndrome with Androgen Insensitivity: Case Report

Androjen İnsensitivitesi Olan Bir Isaacs Sendromu Olgu Sunumu

**ABSTRACT** Isaacs’ syndrome, also known as acquired neuromyotonia, is characterised by hyperexcitability of the motor nerves. This syndrome was first described in 1961 and represents the peripheral nerve hyperexcitability producing muscle stiffness, cramps, fasciculations, myokymia, and pseudomyotonia. In idiopathic forms of neuromyotonia, autoantibodies against voltage gated potassium channels are responsible for the pathogenesis. Secondary forms are mostly related with paraneoplastic conditions. Isaacs’ syndrome tends to associate with some autoimmune diseases. In this case report, we present a 30 year old male patient with muscle twitching and leg cramps due to Isaacs syndrome. Here, the association of polyneuropathy and acquired neuromyotonia occurrence in a patient with androgen insensitivity and vitiligo is discussed.

**Key Words:** Isaacs syndrome; vitiligo; infertility; androgen-insensitivity syndrome


**Anahtar Kelimeler:** Isaacs sendromu; vitiligo; infertilite; androjen duyarsızlığı sendromu

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systemic manifestations like vitiligo and endocrine abnormalities. There have been no previous reports regarding neuromyotonia in association with vitiligo and androgen insensitivity syndrome. As neuromyotonia and several autoimmune diseases including vitiligo are frequently together, we suggest that neuromyotonia may also have an autoimmune association with some endocrine abnormalities.

**CASE REPORT**

In May 2008, a 30-year-old man was referred to our electrophysiology unit for evaluation of leg cramps. Painful muscle cramps and spasms involving both legs and bilateral muscle twitching of the proximal leg muscles were present since childhood. Similar symptoms were present also in his father, but his brothers were asymptomatic. He had impotence and infertility problem that hasn’t been evaluated yet.

Physical examination revealed a continuous, painful muscle spasm with cramps affecting both legs and bilaterally twitching of the quadriceps and semitendinosus muscles. Calf hypertrophy and hyperpigmented skin lesions on lower extremities and abdominal region were also observed. Hyperpigmented areas in his hair were noted.

Neurologic examination showed muscle stiffness and myokymia in gastrocnemius and fasciculations in quadriceps femoris muscles. No muscle weakness was observed. Deep tendon reflexes were decreased. The plantar reflex was flexor. Cerebellar system, cranial nerves and sensory system examinations were normal.

Laboratory blood tests including blood count, serum glucose, blood electrolytes, liver enzymes, creatine kinase, vitamin B12, erythrocyte sedimentation rate, thyroid stimulating hormone, free T3 and T4, anti-T and anti-M antibodies revealed normal results. In addition, anti-nuclear antibody (ANA) and anti-neutrophil cytoplasmic antibodies (ANCA) were negative and anti double stranded DNA (anti-ds DNA) antibody and rheumatoid factor were within normal limits. Tumor markers including carcinoembryonic antigen, alpha-fetoprotein, free prostatic specific antigen, CA 15-3, CA 19-9 and CA 125 were within normal limits. Protein electrophoresis and serum immunoglobulin levels (Ig G, Ig A, Ig M) were within normal limits. Free testosterone, dihydroepiandrosterone and cortisol levels in serum were normal. Total testosterone, adrenocorticotropic hormone, follicle stimulating hormone, luteinizing hormone and prolactine levels were elevated. Serology testing for VGKC antibodies was negative.

Oligoazospermia was noted in sperm analysis whereas abdominal ultrasound and computed tomography (CT) scan of the pelvis were normal. Thyroid Ultrasound, CT scan of the chest and hypophyseal magnetic resonance imaging (MRI) were normal. His electroencephalography testing showed no abnormalities.

Histologic examination of the depigmented area established the absence of melanin using Fontana-Masson stain and the absence of melanocytes in epidermis which demonstrated vitiligo.

Electromyography (EMG) revealed polyphasic motor unit potentials with prolonged latency, short duration and lower amplitudes. Nerve conduction studies showed axonal peripheral neuropathy, the results were shown in Table 1. Conduction block was not detected. Needle EMG showed a few fibrillation potentials and sporadic, spontaneous myokymic discharges in gastrocnemius, anterior tibialis and quadriceps femoris muscles. Several motor units were seen to fire spontaneously and irregularly as doublets, triplets and multiples (Figure 1). Continuous EMG recordings illustrated varying partial motor unit size occuring with high frequency bursts of neuromyotonia in gastrocnemius and anterior tibialis muscles. Caryotype analysis showed 46 XY genotype. Androgen receptor (AR) gene with 26 CAG repeats was noted. IQ test showed mild mental retardation. The patient was treated with lamotrigine (400 mg/day) and carbamazepine therapy (400 mg/day).

**DISCUSSION**

We described the consecutive occurrence of acquired neuromyotonia and mild androgen insensi-
activity syndrome in a patient with idiopathic axonal polyneuropathy and vitiligo. The rare combination of neuromyotonia and androgen insensitivity syndrome has not been reported in the literature before.

We report a patient with acquired neuromyotonia who had muscle stiffness, cramps and myokymias progressing with muscle twitching on proximal lower extremities that exists since childhood. The patient was admitted to our electrophysiology department suffering from myokymias and cramps. Electrophysiologic studies showed decreased recruitment at maximal muscle contraction. Needle EMG showed neuromyotonic discharges in the lower extremity muscles. Our patient’s clinical and electrophysiologic data revealed neuromyotonia with idiopathic subclinical axonal polyneuropathy. Hart reported that nerve conduction studies showed evidence of a polyneuropathy in only one third of patients with acquired neuromyotonia. In most cases in the literature, the neuropathy accompanying neuromyotonia was idiopathic, axonal and subclinical. On the other hand, a number of disorders of peripheral nerve including Guillain-Barre syndrome, chronic inflammatory demyelinating polyneuropathy, hereditary sensory and motor neuropathy have been associated with neuromyotonia. We made the differential diagnosis due to neurological examination, history, clinical and electrophysiologic findings in our patient.

Despite clinical and electrophysiologic evidence of neuromyotonia, antibodies against VGKC were negative. It has been suggested that involvement of the immune system in the pathogenesis of acquired neuromyotonia can not be excluded by negative antibody tests. Moreover, despite of negative testing for VGKC antibodies, acquired neuromyotonia can coexist with axonal neuropathy.

Skin biopsy from hypopigmented lesions demonstrated vitiligo. He also had piabaldism on his hair. Neuromyotonia associated with vitiligo is present in the literature and there are some case reports presented with localized scleroderma.

Our patient has depressed spermatogenesis without any abnormalities in secondary male sex-
ual characteristics. Mild androgen insensitivity syndrome (MAIS) diagnosis was made due to impaired spermatogenesis, male external genitalia on physical examination with normal karyotype analysis, 46 XY and laboratory findings including elevated testosterone and luteinizing hormone levels. Molecular genetic testing for AR gene in blood revealed no mutations. Gottlieb and coworkers reported that somatic mosaism of the AR gene mutations could result in mutations present in DNA in genital skin and not in blood. In addition, they also stated that in some cases with MAIS, only observed abnormality is male infertility.

Although our patient has no mutations in AR gene and presented with neuromyotonia, it is known that Kennedy disease is characterised by CAG repeat expansion in AR gene is accompanying some symptoms of MAIS. Furthermore, CAG repeat expansion may contribute up to 35% of male infertility. We suggest that AR dysfunction which appears to be the only observed abnormality in our patient is leading to infertility. AR gene mutations have been related with some cancers whereas neuromyotonia is related with paraneoplastic conditions and immune mediated disorders. Neuromyotonia with autoimmune etiology is frequently associated with myasthenia gravis, vitiligo, diabetes mellitus, peripheral neuropathy, pernicious anaemia, rheumatoid arthritis and Hashimoto’s thyroiditis. We searched for possible underlying cancers such as lung cancer, thymoma and hematologic malignancies and immune mediated diseases. First, paraneoplastic conditions were investigated. Tumor markers, protein electrophoresis, serum immunoglobulin levels, serum biochemistry, erythrocyte sedimentation rate analysis, abdominal and thoracic CT examinations were found to be all normal. Secondly, our patient had a negative autoantibody screen including anti-thyroid, ANCA, anti-ds DNA, ANA antibodies and rheumatoid factor, plus serum glucose and thyroid function tests were within normal limits.

A few patients with mental retardation and testicular feminization due to AR gene mutations were reported in the literature. There is one report of three brothers with mental retardation who have the shortest CAG repeat in AR gene.

Treatment of neuromyotonia is with anticonvulsant agents such as phenytoin, sodium valproate, lamotrigine and carbamazepine. After we started carbamazepine and lamotrigine in combination, his muscle cramps and spasms improved moderately.

The present case report with acquired neuromyotonia and idiopathic sensorimotor axonal polyneuropathy accompanied with androgen insensitivity syndrome and vitiligo may provide further information about immune system related disorders and endocrine system abnormalities. However, it is still challenging to claim an association between neuromyotonia and endocrine abnormalities without further investigations and studies in the field.

REFERENCES


