Spondyloepiphyseal Dysplasia Tarda with Progressive Arthropathy Associated with Osteoporosis and Cataract: A Case was Misdiagnosed as Juvenile Idiopathic Arthritis

Fatma Gul Yurdakul*, Rezan Kocak, Filiz Sivas, Aysegul Altun Guvenir and Hatice Bodur

Ankara Numune Training and Research Hospital Department of Physical Medicine and Rehabilitation, Turkey

*Corresponding author: Fatma Gul Yurdakul, Ankara Numune Training and Research Hospital Department of Physical Medicine and Rehabilitation Talatpaşa Boulevard No: 5 Altındağ-Ankara, Turkey, Tel: 05059254214, E-mail: fatmagulonder@gmail.com

Abstract

Spondyloepiphyseal dysplasia tarda with progressive arthropathy (SEDT-PA) is a rare hereditary disorder with autosomal recessive, X-linked recessive or autosomal dominant inheritance. Because of joint involvement SED-PA may be confused with juvenile idiopathic arthritis and SEDT-PA also called progressive pseudorheumatoid arthropathy of childhood. The present case had been misdiagnosed as juvenile arthritis and received unnecessary treatments for years.

A 21-year-old male patient admitted with swelling and deformity in the fingers of hands, walking disturbance and common joint pain. Anteroposterior X-ray of the hip joint revealed widened and flattened epiphyses of the femoral head and bilateral narrow and irregular joint spaces. Bilateral tibiofemoral and patellofemoral joint spaces were narrowed in knee radiograph and there was osseous enlargement of the tibiofemoral joints. X-ray of the hands showed periarticular osteoporosis, significant narrowing of the joint spaces of proximal interphalengial and distal interphalangial joints and osseous enlargement of the basis of the metacarpal, proximal and distal phalengeal bones. In lateral vertebral radiograph there was generalized osteopenia, platyspondyly with loss of weight in vertebral bodies, increased kyphosis in lateral thoracic spine and increased lordosis in lateral lumbal spine. He was diagnosed as SEDT-PA with osteoporosis and cortical cataract. Clinical features and joint involvement of SEDT-PA are similar to the JIA. Early diagnosis of SEDT-PA recovers patients from inappropriate and unnecessary treatments. Osteoporosis and ocular anomalies should be considered in patients diagnosed with SEDT-PA.

Keywords
Spondyloepiphyseal dysplasia, Juvenile arthritis, Osteoporosis, Cataract, Misdiagnose

Introduction

The skeletal dysplasias (osteochondrodysplasias) are a group of rare disorders that characterized by common abnormalities in skeleton. The incidence of these disorders is 1/5000. Radiographic abnormalities were evaluated to classify skeletal disorders. In the radiographic classification, these disorders are named according to the pathological area (epiphysyal, metaphysyal, diaphysyal) of the bone. Disorders are named as spondyloepiphyseal dysplasia (SED) when the pathology is in the vertebral epiphysis [1]. SED is defined as 3 clinic groups: 1- SED congenita 2-SED tarda 3-SED tarda with progressive arthropathy (SEDT-PA) [2]. The skeletal dysplasias are genetically heterogeneous and SEDT usually inherited as X-linked recessive. It is characterized by short trunk and extremities, deformity of the vertebrae, platyspondyly, kyphoscoliosis, coxa vara, genu valgum-varum, abnormal shape and structure of the epiphyses of hand bones. Boys are affected more often [1-3]. Although the one set of SEDT-PA is 3 and 6 years of age, they are diagnosed at an older age. Because of joint involvement SEDT-PA may be confused with juvenile idiopathic arthritis and SEDT-PA also called progressive pseudo rheumatoid arthritis of childhood [4-6].

Case Report

A 21-year-old male patient who lives in rural area admitted to outpatient clinic of physical medicine and rehabilitation with swelling and deformity in the fingers of hands, walking disturbance and common joint pain. It started first with low back pain, pain in the hip, knee and ankle joints since 6 years of age. Pain and swelling started in his wrists and metacarpophalangeal joints (PIP) about 10 years old. He had and stiffness lasting for 2 hours. He had been diagnosed as juvenile arthritis. Past medical records were not received. He had received various analgesics and non steroidal anti-inflammatory drugs from different physicians, but he did not remember steroid and DMARD (disease-modifying anti-rheumatic drugs) use. His complaints did not respond to these medical treatments. He had no fever, skin rash, uveitis and muscle weakness. His neuromotor development was normal.

There was consanguineous marriage of his parents. His family members including parents, 4 brothers and 3 sisters had no similar symptoms. His intelligence was normal and had uncomplicated birth history.

Physical examination: His height was 151 cm (vertex to pubis 70 cm and pubis to floor 81 cm), arms span was 166 cm, and weight was 43 kg. Thoracic kyphosis and lumbar lordosis had increased slightly.
distal phalangeal bones (Figure 3). In lateral vertebral radiograph there was generalized osteopenia, platyspondyly with loss of weight in vertebral bodies, increased kyphosis in lateral thoracic spine and increased lordosis in lateral lumbar spine (Figure 4).

Because of generalize osteopenia in vertebral radiographs osteoporosis was assessed with dual energy X-ray absorptiometry (DEXA). L2-L4 bone mineral density was 0.759 g/cm², T-score was -3.0, Femur total bone mineral density was 0.718 g/cm², T-score was -2.4.

He was diagnosed as SEDT-PA with osteoporosis and cortical cataract. He received vitamin D supplementation for Vitamin D deficiency and continued with vitamin D-calcium preparations and antiresorptive treatment. He was advised physiotherapy of joints, informed about his the disease and exercise programs.

Discussion

SEDT-PA is a rare hereditary disorder with autosomal recessive, X- linked recessive or autosomal dominant inheritance [1-3,7]. It is
characterized by irregularities of the end plates of vertebral bodies, swelling and deformities of small and large peripheral joints [6,8]. According to the revision of the International Nomenclature and Classification of the Osteochondrodysplasias SEDTA-PA was listed in the group of “other spondyloepi- (meta)-phyesal dysplasias” [9]. SEDT-PA should be differentiated from juvenile idiopathic arthritis (JIA) to avoid the patients from unnecessary drug use and to ensure the optimal treatment [10]. The absence of laboratory changes indicating systemic or synovial inflammation and lack of response to anti rheumatic drugs help the differential diagnosis of SEDTA-PA from JIA [8,11].

To the best of our knowledge the first case with SEDT-PA was described in 1982. Although the most common initial symptoms of SEDT are low back pain, difficulty in walking and pain in the hips and knees, the disease rarely starts with small peripheral joint involvement. In SEDT patients the age of one set changes between 3-11 years old however the reported age at diagnoses ranges from 4-58 years old [4-20]. The diagnosis is delayed due to the slow progression of symptoms, the rarity of the disease and JIA like joint involvement. Initial symptoms, age of one set and clinical features of some cases in literature are located in table 1 [4-8,10-20]. Our patient had low back pain, pain in hips and knees since 6 years old. Swelling and pain in hand joints started in 10 years old. These features are similar to most of cases we have seen in the literature.

Short stature and longer arm span are known features of the SEDT. Patients’ stature changes between 132-165 cm and arm spans are 152-170 cm [4-7,11,19,20]. The present case height is 151 cm and arm span is 166 cm. He is not taller than the other patients described in the literature. His arm span is longer than his height and upper segment (vertex to pubis) is shorter than lower (pubis to floor) like other patients.

Spine complaints of our patients started in 6 years old and involvement of peripheral joint started in 10 years old. Because of the short stature, affected thoracic and lumbar spine, and hand joints involvement mimicking JIA he was considered as SEDTA-PA. Although clinical findings are similar to JIA, there was no evidence of inflammation in the laboratory findings and radiographic findings were consistent with SEDTA-PA. Autosomal dominant and X-linked inheritance was not considered due to the lack of similar complaints from his patients, sisters and brothers. It is compatible with autosomal recessive inheritance in this patient.

SED'T cases have been published in the literature and disorders have been reported accompanying the SEDTA-PA. Reported disorders include corneal changes and osteoporosis [6,16,19,20]. Although the generalized osteopenia can be seen with SEDT, publications indicate the presence of osteoporosis is limited [13,15,18]. Osteoporosis in SEDT has been reported in 2 cases and both of these cases lumbar L2-4 T score were lower than the hip T score [19,20]. Our patient’s lumbar L2-4 t score was -3.0 and it was lower than hip T score. In addition our patient had also vitamin D deficiency. There were no malabsorption and hydroxylation disorders in this patient. He received insufficient sunlight due to the joint pain and walk disorder. Physician should keep in mind that patients with SEDTA-PA may have osteoporosis and it should be noted that while the treatment.

Another disorder accompanied by SEDT is about cornea and lens. Corneal opacities reported in a patient with SEDT and subcapsular cataract in a patient with SEDTA-PA [6,16]. SEDTA-PA is known as a form of type II collagenopathy and defect in collagen type II may lead to ocular anomalies [16]. In our patient there was bilaterally cortical cataract. The reduced visual acuity in right eye was considered about amblyopia. Patients with SEDTA-PA might have vision problems at an early age and a multidisciplinary approach with ophthalmologist is important.

The treatment of these patients is symptomatic. Growth hormone or anabolic steroid therapy for treatment of short stature was considered but the results were found contradictory. It has been reported in that anabolic steroid therapy can be useful in the treatment of short stature in the patients with skeletal dysplasia in early childhood [21]. On the other hand Burrens’ study results showed that growth hormone therapy is not effective for treatment of short stature in skeletal dysplasia [22]. Specific stretching and strengthening exercises with encouragement for active life are useful therapies for the patients with skeletal dysplasia. Arthroplasty is known to be an option in cases with persistent pain [11]. The present case was advised physiotherapy (Transcutaneous Electrical Nerve Stimulation) for joint pains, informed about his the disease and exercise programs.

In conclusion SEDTA-PA is a rare hereditary arthropathy. Clinical features and joint involvement are similar to the JIA. Early diagnosis of SEDTA-PA recovers patients from inappropriate and unnecessary treatments. Osteoporosis and ocular anomalies should be considered in patients diagnosed with SEDTA-PA.
References


