Case report: Multimodality imaging of van Neck-Odelberg disease

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Abstract

Synchondrosis ischiopubic syndrome (SIS), also known as van Neck–Odelberg disease, is a syndrome characterized by an atypical ossification pattern of the ischiopubic synchondrosis. Its radiological features may mimic stress fracture, neoplasm, osteomyelitis, or posttraumatic osteolysis, causing problems in diagnosis, sometimes leading to unnecessary workup. We report two cases in which the correlation between the clinical and multimodality imaging data enabled the correct diagnosis of SIS.

Key words: Hip pain; ischiopubic synchondrosis; van Neck–Odelberg disease

Introduction

Ischiopubic synchondrosis (IPS) is the junction between the inferior ischial and pubic rami and is principally composed of hyaline cartilage.[1,2] It is a temporary joint, occurring in childhood prior to fusion of the ischial and pubic bones. With skeletal maturation, like all synchondroses, it becomes thinner and then obliterates, either due to bony union or synostosis.[3] IPS closure is somewhat variable and is typically completed before puberty. In early childhood, enlargement of this synchondrosis is bilateral; however, in older children, it is commonly unilateral. Usually, the fusion of the ischial and pubic bones develops without any clinical symptoms. However, sometimes children may have pain in the hip, in the groin, or in the gluteal region, resulting in limitation in the movement of the hip joint and limping.[4,6] These clinical features of the synchondrosis ischiopubic syndrome (SIS) are nonspecific and create problems in the differential diagnosis of equivocal findings on plain radiographs. Odelberg (1923) and van Neck (1924) described radiographic changes of SIS as swelling and demineralization of the ischiopubic fusion zone and referred to this entity as “osteochondritis ischiopubica.”[9,10] We report here two children with acute hemipelvis pain who were correctly diagnosed to have SIS.

Case Reports

Case 1

An 8-year-old boy with left foot dominance presented with acute pain in the right ischiopubic region, with a limp for 1 month. There was no history of trauma. There was no palpable swelling of the IPS, and contractures of the adductor muscles were not observed with either rectal or external palpation. He had never been treated with antibiotics. Laboratory tests revealed a slight increase in the C-reactive protein (CRP) and the erythrocyte sedimentation rate (ESR). Anti-streptolysin-O (ASO) titer was normal. Anteroposterior radiographs of the pelvis showed an enlarged right IPS characterized by focal areas of osteolysis and associated sclerosis and irregular edges [Figure 1 A,B]. A CT scan study, using a 64-detector row unit (Aquilion 64, Toshiba Medical Systems, Tokyo, Japan) better defined the areas of demineralization at the edges of the synchondrosis with associated sclerosis and irregular edges [Figure 3]. A remarkable improvement in symptoms was observed after 3 weeks of anti-inflammatory therapy and bed rest. One
month later, a second MRI showed a decrease in perilesional edema [Figure 4].

Case 2
A 12-year-old right-footed male presented with left hip pain and inability to walk. He denied traumatic events and reported difficulty in playing football. Moreover, he had never been treated with antibiotics or anti-inflammatory drugs. Laboratory tests were unremarkable except for a slight increase in CRP. ASO titer was also normal. Radiographs of the pelvis showed rarefaction of the left IPS with irregular and puffy bone edges [Figure 5]. A CT scan demonstrated diastasis with sclerotic borders [Figure 6]. MRI revealed fusiform enlargement of the left IPS without any soft tissue involvement [Figure 7]. After anti-inflammatory treatment and bed rest for 2 weeks, there was complete regression of symptoms.

Discussion

A radiolucent swelling at the ischiopubic fusion zone in the prepubescent skeleton was firstly described by Odelberg (1923) and Van Neck (1924) as “osteochondritis ischiopubica.”[9,10] At present, this finding is considered a normal ossification pattern[11] since closure of the IPS is variable and can occur at an age from 4 to 16 years. Asymmetry in closure is the rule rather than the exception. In fact, in older children, before complete ossification, a unilateral enlarged IPS is commonly seen.[12] As reported in literature, unbalanced mechanical stress, such as kicking or jumping, may induce an inflammatory reaction and a delayed complete ossification of this synchondrosis.[2,3,13,14] The IPS is composed of hyaline cartilage; and its cell layers, before fusion, show a strong enhancement on hematoxylin
and eosin staining. This is typical of joints subjected to mechanical stress.

SIS involves hip pain, sometimes involving the gluteal region, causing marked limping and spontaneous tension of the adductors not related to trauma. In fact, in our two patients, we observed a slight limp and limitation in flexion and extension. In both the boys, the side of IPS correlated with foot dominance,[15] which probably may suggest that SIS is a physiologic reaction to forces exerted on the nondominant limb during physical activity.

Its tumor-like appearance on conventional radiographs may be mistaken for stress fracture, tumor, or inflammation. The radiographs and CT scans are usually typical showing increased size with lucencies and sclerosis, as seen in our patients. The typical MRI features such as signal alteration of the bone marrow and fusiform swelling of the adjacent soft tissue were also seen in our patients. We also detected a hypointense band-like structure perpendicular to the pubic axis, defined by Herneth et al. as “fibrous bridging,” a typical MRI feature of this structure.[2,16,17]

Differential diagnosis can be difficult with pubic rami stress fractures, which are very common lesions and occur in athletes, or after radiation therapy.[2,16,17] A stress fracture typically presents with hyperintense marrow edema on T2W images and a hypointense irregular fracture line perpendicular to the long axis of the superior pubic ramus. Tumors such as Ewing’s sarcoma usually present with permeative bone destruction and extension into the soft tissues. However, according to Kloiber et al.,[18] initial radiographs were useful as a baseline study to identify further demineralization and destruction that occurred in osteomyelitis. In osteomyelitis, bone destruction and abnormal soft tissue are seen[18] with peripheral enhancement and abscess formation.[19]

Knowledge of this condition is essential to make sure that it is not mistaken for stress fracture, infection, or tumor, in symptomatic children.

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