Osteoarthritis at Young Age, a Diagnostic Challenge: A Case of Stickler Syndrome

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Abstract: A young woman presents with severe polyarticular osteoarthritis with relevant family history potentially suggesting a hereditary disease. Previously, the patient’s mother had been diagnosed with rheumatoid arthritis and reported to have suffered from some locomotor problems. Careful clinical evaluation with an extensive personal and familial history pointed towards a diagnosis of Stickler syndrome, an autosomal dominant condition with progressive arthro-ophthalmopathy, (early osteoarthritis and myopia). Following this timely diagnosis, genetic counselling was offered.

Keywords: Myopia, osteoarthritis, scoliosis, Stickler Syndrome.

INTRODUCTION

Complete and timely medical diagnoses rely on the identification of symptom patterns that are characteristic for certain patient demographics. Therefore, it is plausible that a disease presenting at an unusual age might be overlooked as part of the differential diagnosis. Furthermore, a disease commonly associated with the elderly that presents at a younger age might indicate a hereditary disease; in which a diagnosis is a priority for genetic counselling. Here, we report a patient presenting with early onset polyarticular osteoarthritis, with a potential hereditary cause. A full clinical evaluation and an extensive personal and familial history taking resulted in the correct diagnosis. The patient gave oral informed consent for this report.

CASE REPORT

A 26-year old female patient presents with arthralgia and intermittent swelling of the PIP joints, ankles and knees of several years. Symptoms worsen on joint loading and when participating in sports, there was no morning stiffness or pain at night. A brief medical history shows that she underwent a fusion procedure for scoliosis and a Ravitch procedure for pectus carinatum during childhood. She suffered a hip fracture aged 18 and was previously diagnosed with myopia. Clinical examination revealed a woman with short stature (1.60m) with exceptionally long and thin fingers, a status post scoliosis fusion and chest surgery, and mild hydrops with flexion deficit in the right knee. Applying patellar pressure was painful for both knees. Radiology revealed significant degenerative changes of the knees, ankles and some PIP joints (Figs. 1, 2).

Fig. (1). X-rays showing the patient’s unusually slender hands and long phalanges (arrow), a feature that can be seen in patients with Stickler syndrome.

The patient was accompanied to the first outpatient visit by her mother, who had previously undergone a bilateral hip replacement in her thirties. The patient’s mother was also treated for presumed rheumatoid factor negative rheumatoid arthritis (RA). The patient’s grandfather also suffered from “early arthrosis”. A new radiological evaluation was requested from the rheumatologist overseeing the mother’s follow up care. Results from this clearly showed polyarticular osteoarthritis with no typical RA erosions (Figs. 3, 4).

The main reason for this young female to attend a consultation was her concern about an inheritable disease. The combination of familial osteoarthritis at a young age, early scoliosis and the myopia suggests a diagnosis of autosomal dominant Stickler syndrome; in depth analysis revealed a c.2155C>T mutation in the COL2A1 gene corresponding with Stickler syndrome.
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DISCUSSION

Gunnar B. Stickler first described Stickler syndrome in 1965 as an autosomal dominant progressive arthro-ophthalmopathy [1]. Disease manifestations are variable with a late or misdiagnosis being common. Clinical features consist of orofacial and ophthalmic features, deafness and osteoarthritis. Typical orofacial features include a flat midface with a depressed nasal bridge, anteverted nares and micrognathia [1-4]. The majority of families with Stickler syndrome have mutations in the COL2A1 gene: Type 1 or membranous-vitreous type. Others may have mutations in the COL11A1 gene, known as Type 2. Mutations in COL11A2 have also been described, displaying a more systemic syndrome but no ophthalmological features, known as Type 3. Other loci are yet to be identified.

Osteoarticular manifestations are common and include early-onset osteoarthritis, spinal abnormalities and joint hypermobility, with up to 50% of patients developing osteoarthritis before the age of 30 [5]. The typical orofacial features were not present in this patient, although these can become less expressive with increasing age, and the patient suffered no hearing loss. The slender fingers seen here are infrequently reported in literature. In summary, our patient was suffering from early osteoarthritis in the knee joints, like her mother who had previously undergone replacement knee surgery. There was no valgus component which is often reported. As osteoarthritis in Stickler syndrome is more frequently seen in weight-bearing joints like the hips or knee, this, coupled to the other skeletal features of a pectus carinatum and scoliosis and the myopia were compatible with a diagnosis of Stickler syndrome [6].

This case report highlights that detailed personal and familial history and full clinical evaluation are crucial to making diagnosis in patients with unusual symptom presentation. The importance of timely diagnoses for hereditary disorders, cannot be emphasised enough to ensure adequate genetic counselling.

Furthermore, when reviewing the patient’s file from the orthopaedic department at the time of scoliosis surgery, it was discovered that a diagnosis of Stickler syndrome had been suggested, although no diagnostic confirmation was carried out. Despite this, the patient, her mother nor the patient’s GP were aware of this syndrome. This highlights the importance of proper transition of youngsters with serious diseases from paediatric into adult care.

ABBREVIATIONS
DIP = Distal Interphalangeal
GP = General Practitioner
MCP = Metacarpophalangeal
PIP = Proximal Interphalangeal
RA = Rheumatoid Arthritis

CONFLICT OF INTEREST

The authors confirm that this article content has no conflict of interest.
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