Unusual Presenting Syndrome of Rheumatoid Arthritis Exacerbation

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ABSTRACT

We herewith report the rare case of a patient with juvenile rheumatoid arthritis who developed on 2 occasions the clinical picture of an isolated unilateral palsy of the inferior oblique muscle—Brown’s syndrome, following pregnancies, each time in a different eye. Although the eye is frequently involved in rheumatoid arthritis (RA), the Brown’s syndrome is seldom reported in literature with regards to RA.

Keywords: Brown’s Syndrome; Juvenile Rheumatoid Arthritis; Rheumatoid Arthritis

1. Introduction

The eye is frequently involved in rheumatoid arthritis (RA). The most common manifestations are keratoconjunctivitis sicca, often resulting in corneal involvement, scleritis and episcleritis. However, other less frequent ocular features have been described [1]. We herewith report the case of a patient with juvenile rheumatoid arthritis who developed on 2 occasions the clinical picture of an isolated unilateral palsy of the inferior oblique muscle. Following pregnancies, each time in a different eye.

2. Case Report

A 40 years old woman, with a 25 year history of juvenile rheumatoid arthritis (JRA), developed sudden right eye gaze palsy together with arthritis exacerbation the first week after giving birth to her second child. She reported a normal 1st pregnancy without after delivery complications. At this time, a few days after completing her second pregnancy, she presented with complains of diplopia at an upward-in gaze. During the pregnancy, the patient entered into remission of JRA, without need for medications. On physical examination, at presentation, she demonstrated severe synovitis of wrists, 2, 3, MCPs, 2 - 4 PIPs and knees. The ophthalmologic examination was consistent with right inferior oblique palsy. Laboratory tests revealed a normocytic-normochromic anemia, and raised ESR and CRP. Single bundle EMG of the ocular nerves excluded neurological pathology and muscle myopathy.

The patient was diagnosed as suffering from Brown syndrome. She was treated with Prednisolone 30 mg a day and Methotrexate (MTX) 15 mg a week. Her diplopia slowly improved during the course of several weeks, resulting in a slight permanent disturbance of eye movement.

A few years later, she had an uneventful 3rd pregnancy but after her fourth delivery, she presented with a clinical picture similar to the one observed after her second pregnancy: arthritis exacerbation and diplopia. This time, the event involved the left eye, at an upward-in gaze. The patient was examined by an ophthalmologist, and diagnosed again as Brown syndrome. Treatment with MTX 15 mg a week was renewed and Prednisolone dose raised to 30 mg a day (the patient was treated by prednisolone 5 mg once a day during her fourth pregnancy). Her eye symptoms improved, although a mild residual diplopia remained.

The patient was treated by Prednisone 30 mg/d till the beginning of improvement in her diplopia with slow Prednisone tapering after that in both cases.

3. Discussion

In 1973, Brown reported an unusual motility disorder, characterized by a limitation of upgaze in an adducted eye [2]. The cause of this malfunction is an abnormality of the superior oblique tendon, its sheath or the trochlea, preventing free passage of the tendon through the trochlea during inferior oblique action. These clinical features have been reported in association with SLE [3],
Sjogren’s syndrome [4], JRA [5] and RA [6]. In the latter, a stenosing tenosynovitis of the superior oblique tendon sheath causes mechanical obstruction when attempting to raise the eye in adduction. The problem resembles trigger finger in the hands of RA patients [7]. Despite the common association of tenosynovitis with rheumatoid arthritis, Brown syndrome is a rare complication and has been seldom reported. While reviewing the literature, we found only sporadic cases describing Brown’s syndrome in association with RA or JRA. Hickling and Beck [8] reported 6 cases during a period of 10 years. Other case reports describe only one or two patient in each case. Interestingly, Brown’s syndrome may appear when RA is quiet under DMARDs treatment [9], as well as during disease exacerbation. On the other hand, spontaneous resolution of this condition may occur [10]. The traditional treatment of Brown’s syndrome is corticosteroids with complete resolution of the disease in most cases [11]. Our case is also unique since it appeared after pregnancy as the first presenting sign of RA exacerbation and recurred in the same circumstances in the another eye. The mild residual symptoms observed in our patient may suggest a possible congenital malformation of the tendon.

4. Conclusion

We have reported a rare case of recurrent Brown syndrome after pregnancy in a patient with JRA. Because of rarity of this syndrome, it is probably often under diagnosed. Increased awareness to the possibility of tenosynovitis of the superior oblique tendon as a manifestation of active RA may prompt its diagnosis and treatment.

REFERENCES


