Magnetic Resonance Imaging Findings of Elbow Involvement in Behçet’s Disease: Medical Education

Behçet Hastalığında Dirsek Eklemi Tutulumunun Manyetik Rezonans Görüntüleme Bulguları

ABSTRACT Behçet’s Disease is a systemic vasculitis of unknown etiology. Articular involvement is a well-recognized feature, that is generally intermittent, self-limiting, non-erosive, non-destructive, and non-deforming, usually monoarticular or symmetrically oligo-articular subacute arthritis. While knees, ankles and wrists are most commonly affected, elbow involvement is very rare. Previous descriptions of articular involvement have been mostly limited to conventional radiographic findings. We report magnetic resonance imaging (MRI) findings of pathologically proven acute synovitis on chronic arthritis of the left elbow in a 25 year-old man with Behçet’s disease, who followed up for 9 years.

Key Words: Magnetic resonance imaging; Behcet Syndrome; arthritis


Anahtar Kelimeler: Manyetik rezonans görüntüleme; Behçet hastalığı; kronik artrit


Behçet’s disease, which is a systemic vasculitis, may commonly exhibit articular involvement; however the elbow is very rarely affected. Plain radiographs and clinical findings may not be satisfactory to diagnose the degree of involvement. MRI shows clearly the presence of synovial proliferation and inflammation. MRI findings in the case with Behçet’s disease demonstrating acute synovitis in chronic arthritis of the left elbow are discussed in this paper.

A 25-year-old man was presented with progressive left elbow pain, which had started about 9 months ago. The patient had a 9-year history of recurrent oral and genital ulcerations, uveitis, and papulopustular skin lesions. He had been diagnosed with Behçet’s disease; however, the follow-up was irregular. Physical examination showed swelling and tenderness of
the left elbow as well as limited range of motion. Plain radiograph showed joint-space narrowing, marginal erosions along the radial side of the elbow joint and soft tissue swelling (Figure 1). Informed consent was obtained. MRI was performed via 1.5-T MR scanner (Philips Gyroscan Intera Master, Eindhoven, The Netherlands, 30 mT/m maximum gradient strength, 150 mT/m per millisecond slew rate) using a surface coil. The patient was placed in the prone position with his left arm extending next to his head. MRI protocol included T1 (TR/TE=578/15, TSE factor: 3, matrix: 320 x 512, NSA: 4) and T2 (TR/TE=2500/100, TSE factor: 21, matrix: 320 x 512, NSA: 3) weighted TSE sequences; additional fat saturated T2 weighted images were performed in axial, sagittal and coronal planes. MRI revealed joint effusion and synovial thickening with marginal erosions and soft tissue edema (Figure 2a, 2b). After intravenous administration of 0.1 mmol/kg contrast medium (gadolinium-DTPA, Magnevist®, Schering, Germany), there was enhancement of the inflamed synovium due to acute synovitis (Figure 2c). The patient was diagnosed

FIGURE 1: Plain radiograph shows joint-space narrowing, marginal erosions along the radial side of the elbow joint with soft tissue swelling.

FIGURE 2: Sagittal A. precontrast T1-weighted, and B. T2-weighted images demonstrate joint effusion, significant synovial hypertrophy and soft tissue edema. C. Marked enhancement of the inflamed synovium following intravenous gadolinium is seen on post-contrast T1-weighted image.
with arthritis caused by Behçet’s disease. He was given indomethacin 125 mg/day, prednisolone 10 mg/day and sulphasalazine 2 g/day. The findings did not change significantly on follow-up MRI examination at 3 months after treatment (Figure 3). Although intraarticular corticosteroid injection was performed, no clinical response was achieved notably, so a synovial biopsy was carried out to rule out any infectious disease including tuberculosis. Since pathological examination showed non-specific acute synovitis on chronic base, azathioprine 125 mg/day was added to the treatment (Figure 4a and 4b).

**DISCUSSION**

Behçet’s disease is a systemic vasculitis of unknown etiology with the classical triad of oral and genital ulcerations and uveitis. The diagnosis is based on the criteria of the International Study Group defined in 1990. Ocular, articular, vascular, pulmonary, mucocutaneous, gastrointestinal and nervous tissues are prone to develop manifestations of Behçet’s disease. Although articular involvement is a well-recognized feature (5% to 97% prevalence), the elbow is very rarely affected. Children are more likely to present with joint manifestations (73.7%) and polyarthritis (35.7%) than adults. Articular involvement is generally described as an intermittent, self-limiting, non-erosive, non-destructive, and non-deforming, usually

**FIGURE 3:** Sagittal T2 weighted images on the follow-up period at 3 months demonstrate significant synovial proliferation.

**FIGURE 4:** A. Villus like projections of hypertrophied synovium with diffusely proliferating mononuclear inflammatory cells (H&E x40). B. Acutely inflamed fibrinous surface of the synovium with underlying vascular proliferation and chronic inflammation (H&E x40).
monoarticular or symmetrically oligo-articular subacute arthritis. Knees, ankles and wrists are the most commonly involved joints, while involvement of the spine, sacroiliac joints, shoulder, hip are rare and elbow is very rare.²,⁵⁻⁹ The duration of attacks is usually 2 months or less, but may occasionally be as long as 4 years.²,⁵⁻⁸ Unusual forms include polyarthritis resulting in joint deformity and/or destruction.⁵,¹⁰,¹¹

Imaging findings include mostly non-specific findings such as osteoporosis and soft-tissue swelling, rarely accompanied by joint-space narrowing and marginal erosions.³⁻⁵,¹¹,¹² Enhancement due to gadolinium uptake in the synovium may correlate with acute synovitis. Biopsy may confirm the imaging findings of synovial proliferation and inflammation.⁶,¹²⁻¹⁴ Synovial inflammation was documented in previous reports. It is confirmed by synovial fluid, usually a good mucin clot formation, hypertrophy and hyperplasia of synovial lining cells, hypervascularity, subsynovial accumulation of inflammatory cells, and replacement of the superficial zones of the synovial membrane by dense inflamed granulation tissue composed of lymphocytes, macrophages, fibroblasts, neutrophils, and vascular elements.⁵,¹²,¹⁴ Rheumatoid arthritis is the main entity in the radiological and pathological differential diagnosis.¹⁴

Our case demonstrated monoarticular, chronic, erosive, deforming elbow involvement. The cardinal radiological findings consisted of joint-space narrowing, marginal erosions along the radial side of the elbow joint accompanying soft tissue swelling and acute synovitis with synovial proliferation and inflammation. Long-standing Behçet’s disease, monoarticular involvement, negative rheumatoid factor and typical MRI findings lead us to diagnosis of arthritis due to Behçet’s disease. There was no clinical response to non-steroidal anti-inflammatory drugs, oral and intraarticular corticosteroid treatment and sulphasalazine. Thus biopsy was performed. Histologically, synovial tissue showed villus like hypertrophy with prominent inflammation, in part, synovial surface had fibrinous material and intimately mixed polymorphonuclear leukocytes were seen. In other areas and beneath this acutely inflamed surface there was vascular proliferation and prominent, vaguely nodular lymphoplasmocytic infiltration in the synovial tissue. There were neither giant cells nor granulomatous stuctures detected. The patient is currently taking azathioprine in addition to his previous medications.

CONCLUSION

Although very rare, Behçet’s arthritis should be considered for the differential diagnosis of elbow arthritis. MRI gives information about the degree and phase of synovial involvement and is helpful for the evaluation of treatment.

REFERENCES