Isolated digital vasculitis in a patient with rheumatoid arthritis: good response to tumour necrosis factor $\alpha$ blocking treatment

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Abstract

Tumour necrosis factor $\alpha$ (TNF$\alpha$) blocking agents are among the most promising new treatments for rheumatoid arthritis (RA). However, no data exist about the effect of these agents on extra-articular manifestations of RA. A patient is described with small vessel vasculitis that repeatedly responded well to treatment with the soluble p55 TNF$\alpha$ receptor fusion protein Ro 45-2081 (lenercept).

Symmetrical polyarthritis is the hallmark of rheumatoid arthritis (RA). In addition, pathology outside the joints can also be found. One of these so-called extra articular manifestations of RA is vasculitis. Vasculitis in RA usually affects small vessels and commonly involves the skin, causing nailfold infarcts, and in more severe cases digital gangrene and leg ulcers. Less frequently, vasculitis of small and medium sized arteries complicates RA and causes damage to peripheral nerves and internal organs. Prevalence of vasculitis in RA varies from 1 to 5% but postmortem studies show higher percentages up to 25%. Treatment of rheumatoid vasculitis depends on the size of the vessels affected and the impending organ damage. High doses of corticosteroids may be necessary, often in combination with immunosuppressive agents, including azathioprine and cyclophosphamide, to allow for long term disease control and reduction of the corticosteroid dose. Treatment with these drugs is associated with considerable side effects.

In the past decade new agents have been developed for the treatment of autoimmune diseases. Among the most promising are tumour necrosis factor $\alpha$ (TNF$\alpha$) blocking agents. Several agents that block TNF$\alpha$ have been developed and studied in patients with RA. TNF$\alpha$ blockade is achieved either with monoclonal antibodies against TNF$\alpha$ or with fusion proteins containing human TNF receptors bound to an Fc component of a human IgG antibody. Although excellent efficacy in treatment of RA has been reported for several of these agents, no reports have thus far considered the effect of TNF$\alpha$ scavenging on extra-articular manifestations. We describe a chance observation in a patient with RA and nailfold lesions, responding repeatedly to treatment with the anti-TNF$\alpha$ receptor fusion protein, lenercept.

Case report

A 46 year old woman was diagnosed with rheumatoid factor positive, erosive RA in 1982.
From the beginning of her disease, antinuclear antibodies could be detected. Testing for disease-specific autoantibodies was negative on several occasions. Her medical history was unremarkable. In the following years she was treated unsuccessfully with hydroxychloroquine, intramuscular gold salts, d-penicillamine, azathioprine, methotrexate, and a combination of sulfasalazine and methotrexate. Because of progressive joint destruction she received shoulder prostheses on both sides, an elbow endoprosthesis left, total knee joint replacement on both sides, and a spondylosis of the first and second cervical vertebra was performed. No extra-articular symptoms were present except for sicca complaints. Owing to the uncontrollable disease she was included in 1994 in a study with Ro 45–2081, a fusion protein combining two p55 TNF receptors with the Fc component of an IgG human antibody (lenercept). After a three month placebo controlled phase she was treated with 50 mg lenercept intravenously every four weeks. Clinical response was impressive with swollen joint counts decreasing from 32 to five and C reactive protein (CRP) levels declining from 95 mg/l at baseline to 20 mg/l after the first injection. Low disease activity was sustained for the following years. Besides lenercept, her drug treatment consisted of oral prednisone 5 mg a day and, occasionally, paracetamol 500 mg.

In the spring of 1999 she first noticed nailfold lesions on the fingers of both hands. These lesions disappeared after every injection of lenercept and reappeared three weeks thereafter when the effect of lenercept was decreasing (figs 1A and B). Clinical response indicated by a drop in joint counts and CRP was present two weeks after each anti-TNFα injection.

Discussion
We describe a patient with RA and small vessel vasculitis manifested by nailfold lesions responding to treatment with lenercept. The response has been well documented during two cycles of anti-TNFα administration. Although