Letters

NO DETECTION OF MACROPHAGE ERYTHROPOIETIN PRODUCTION IN BONE MARROW FROM RHEUMATOID ARTHRITIS PATIENTS WITH AND WITHOUT ANAEMIA AND CONTROLS

To the Editor

Anaemia is a common extraarticular manifestation of rheumatoid arthritis (RA). Apart from deficiencies of iron, vitamin B12 and folic acid and adverse antirheumatic drug reactions, the anaemia of chronic disease (ACD) is the most frequent cause of anaemia in RA (1). Much research has been carried out to examine its pathophysiology, such as altered iron metabolism (2,3) and cytokines — like interleukin 1 and tumour necrosis factor, which may suppress erythropoiesis (4). Another mechanism underlying ACD in RA might be an insufficient erythropoietin (Epo) responsiveness to the anaemia (1,5) or decreased erythroblast-Epo sensitivity based on findings that Epo-induced erythroid growth is reduced in RA complicated by ACD (6). Rich reported that macrophages in bone marrow are able to produce and release Epo depending on local pH and oxygen tension (7) and exhibit Epo gene expression (8). These observations prompted us to investigate whether Epo production is detectable in bone marrow derived from RA patients and controls and whether Epo production by marrow macrophages and its metabolism is altered in ACD.

Serum and bone marrow were obtained from 5 nonanaemic RA patients, 5 RA patients with ACD (based on normal to increased bone marrow iron stores and the exclusion of other causes of anaemia) and 4 controls (bone marrow transplant donors), after they gave written informed consent. Mean age of the RA patients was 59(±6) years and of the controls 32(±4) years. None of the RA subjects were treated with steroids or cytostatic drugs. After Ficoll separation, one million mononuclear cells were added to a medium containing the usual substances for BFUe colony assessment (9), up to volume of 1 ml. It was then incubated for 48 hours at 37 degrees centigrade and 100% humidity in 5% CO2. This procedure was repeated with the addition of 1 U/ml of Epo (Boehringer Vienna, Austria). The cell suspension was centrifuged and the supernatant was tested for Epo, using an ELISA Epo assay as recently developed (Boehringer Mannheim, W-Germany; lowest detection level 0.1 mU/ml; reference values in serum 0.4-8.9 mU/ml). The results are shown in Table I. No Epo could be detected in any of the marrow cultures of the three groups. After the addition of Epo to the cultures less than 50% was recovered after 48 hours of culturing.

These observations may indicate that Epo production and/or release by marrow macrophages in culture is not detectable in the three groups studied here. The second experiment shows that recovery of Epo added to the cultures is less than 50%, suggesting that Epo is metabolized, adhered to or incorporated into the marrow cells, which were washed away in these experiments. The Epo initially produced in the marrow cultures in the first experiment might have been below the detection limit or metabolized before measurement. Apparently net macrophage Epo production or release after 48 hours of culturing is undetectable in either group and Epo metabolism is similar in the three groups. Consequently, these preliminary data do not support the concept of altered marrow Epo production or metabolism by macrophages or other cells present in bone marrow as a cause of ACD in RA.

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REFERENCES


HERPES SIMPLEX VIRUS ANTIGENS IN SERA OF PATIENTS WITH BEHÇET'S DISEASE

To the Editor:

In a recent study, it was reported that titres of IgG antibodies to herpes simplex virus type 1 (HSV-1) were significantly higher in the sera of patients with Behçet's disease (BD). Levels of IgG antibodies were significantly higher in patients with active disease than in patients with inactive disease. These results suggested some evidence for involvement of HSV-1 in the immunopathogenesis of BD (1). In the present study we investigated HSV antigens in the sera of patients with BD. Twenty patients (19 male, one female) with a mean age of 39 ± 3 (range 29-58) were studied. Disease duration was 6.8 ± 1.9 (range 2-25). All patients fulfilled the diagnosis criteria proposed by the International Study Group for Behçet's disease (2). Clinical manifestations were as follows: buccal aphthosis (n = 20) genital aphthosis (n = 17), necrotic pseudofolliculitis (n = 15), dermohypodermal lesions (n = 5) positive pathergic test (n = 13) venous thrombosis (n = 8), arthritis (n = 7) central nervous system (= 3) arterial aneurysm (n = 2). The disease was active in 12 patients and inactive in 8. Antigens present in sera of BD patients at different stages were investigated using an ELISA technique. Monoclonal antibodies against HSV-1 and 2 were fixed on wells of polystyrene plates. 100 microliters of sera were incubated pure and at 1:10 for one hour at 37°C. After washing, 100 microliters of anti HSV monoclonal antibody was added (1:50 for one hour). After washing, phosphatase alcaline conjugate was added for 1 hour (100 microliters). Finally, 100 microliters of phosphatase alcaline substrate were used for 20 min. Positive controls used were HSV-1 and 2 strains purified in ultracentrifugation gradient sucrose and ultrasonicated. Negative controls were PBS. Positive controls had a high optical density at 405 nm (>0.6). Negative controls were always <0.15.

On the 20 patients tested, none had positive results by this assay. Reactivation of the BD does not seem to be associated with liberation of HSV particles in the sera but with rise of antibodies against HSV as previously described.

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