CASE REPORTS

Megaloblastic anaemia due to trimethoprim-sulphamethoxazole therapy in uraemia

G. M. YUILL
M.B., Ch.B., M.R.C.P., B.Sc.

University Hospital of South Manchester

Summary
A patient is described in whom malignant hypertension led to oliguric renal failure. A course of trimethoprim and sulphamethoxazole (co-trimoxazole) was prescribed for an intercurrent urinary tract infection and appeared to precipitate a megaloblastic anaemia which responded satisfactorily to folic acid.

Introduction
Sulphonamides interfere with the conversion of para-aminobenzoic acid to dihydrofolic acid. Trimethoprim interferes with the conversion of dihydrofollic acid to tetrahydrofollic acid by interfering with the enzyme dihydrofollic acid reductase. The bacterial enzyme is \( \times 10,000 \) more sensitive than the mammalian enzyme to trimethoprim. Usually this selective toxicity enables one to eradicate bacterial infection without significantly interfering with the metabolism of the mammalian host. However, there are several reports demonstrating an effect of the trimethoprim–sulphonamide combination on human folic acid metabolism.

Kahn, Fein & Brodsky (1968) administered trimethoprim to a group of patients with normal blood counts and blood urea. One patient's haemoglobin fell from 14:0 to 8:7 g/100 ml; another's platelet count fell from 200,000 to 60,000/mm\(^3\). They also observed a rise in the polymorphonuclear lobe count, a rise in FIGLU excretion, and giant metamyelocytes and transitional megaloblastic erythroid changes in the bone marrow.

They treated a second group of patients with trimethoprim and sulphamethoxazole (co-trimoxazole) and reported the following abnormalities: leucopenia, granulocytopenia, thrombocytopenia, and an increase in FIGLU excretion. Bone marrows were not examined in this group.

Allison et al. (1969) and Hulme & Reeves (1971) also reported similar abnormalities.

The current case report is of a patient who was given co-trimoxazole whilst oliguric and who developed severe megaloblastic anaemia, thrombocytopenia and mild leucopenia.

Case history
J.B. was a 40-year-old housewife who was admitted to Withington Hospital in June 1971. Three months before admission she developed breathlessness on exertion and general malaise. She was discovered to have severe hypertension which proved difficult to control and her renal function had deteriorated steadily. She was transferred to Withington Hospital.

On examination she was pale, breathless and anxious. BP 200/125 mmHg; CVP elevated to 6 cm H\(_2\)O. There were crepitations at both lung bases, and oedema of the face, sacrum and ankles. She had a grade 4 hypertensive retinopathy. She had psoriatic patches over the trunk and her scalp and a few psoriatic pits on the nails. There was thickening of the metacarpophalangeal joint of the left thumb and the left little finger and there was slight thickening of the periarticular tissue around the knees. A clinical diagnosis was made of essential hypertension entering a malignant phase with renal failure. In addition she had a mild inactive psoriatic arthropathy.

Investigations. Hb 6·5 g/100 ml, WCC 12,000/mm\(^3\), platelets 40,000/mm\(^3\), reticulocytes 9%, blood urea 254 mg/100 ml; sodium 142 mEq/l, potassium 4·6 mEq/l, chloride 90 mEq/l, urea clearance < 1 ml/min. Urine volume about 200 ml/day. Serum albumin 3·5 g/100 ml, globulin 3·1 g/100 ml, bilirubin, thymol turbidity, alkaline phosphatase, transaminases and serum protein electrophoresis normal. Calcium 9·8 mg/100 ml, phosphorus 7·4 mg/100 ml. Serological tests for SCAT, latex, antinuclear factor, LE cells, WR, Coombs' test were negative. The ASOT was normal. The kidneys were 11 and 11·5 cm in length. The electrocardiograph showed left ventricular hypertrophy and strain and left axis deviation.

Progress. She was put on an 18 g protein, 20 mEq sodium Giovanetti diet with vitamin and methionine supplements. Blood pressure was controlled with methyldopa 250 mg q.d.s. and she had several peritoneal dialyses in order to remove excess fluid. Investigations revealed the presence of a significant
growth of coliforms sensitive to Seprin in the urine on four occasions and on 25 August she was put on tabs. Seprin 1 q.d.s. (sulphamethoxazole 1·6 g and trimethoprim 320 mg daily) in addition to the above regimen. She was treated with Seprin until 14 September when the urine became sterile. During this period she was found to have a moderate reticulocytosis, a low platelet count, and significantly raised fibrin degeneration products. The haemolytic uraemic syndrome was diagnosed with significant intravascular coagulation as a complication of malignant hypertension and in view of her serious clinical state, on 9 September, she was started on intravenous heparin therapy which continued until 15 September. During this time it was noticed that her reticulocyte count and platelet count were falling.

The falling platelet count was attributed to haemolytic uraemic syndrome, but the fall in the reticulocyte count was not explained. At the same time it was noticed that the haemoglobin was falling steadily despite no overt source of blood loss and reasonable control of her uraemia. A blood film at this time showed some hypersegmented polymorphs, and occasional macrocytes. The platelet count reached 40,000 and a sternal marrow was performed which demonstrated megaloblastic changes in the erythroid series and occasional giant megalocytes and raised iron stores. She was treated empirically with 1000 μg of vitamin B₁₂ daily, 15 mg of folic acid parenterally daily from 10 September for 1 week and the Seprin was stopped on 13 September. On 16 September her haemoglobin was rising and she had a brisk reticulocytosis and the platelet count had reached 200,000. The serum B₁₂ + folate analysis was then returned, the B₁₂ was 295 pg/ml and the serum folate level was 0·1 ng/ml and further B₁₂ therapy was stopped (Fig. 1).

She was established on intermittent haemodialysis on 2 October, with the blood urea ranging from 60 to 80 mg/100 ml and by 12 October the haemoglobin was 6·9 gm/100 ml, white cells 9000/mm³, and reticulocytes 6%. A repeat sternal marrow was performed (15 October) and showed normoblastic erythropoiesis with strikingly deficient iron stores, a finding in marked contrast to the previous marrow. There was defective haemoglobinization and she is currently being treated with a course of intravenous iron. On 23 November her haemoglobin was 11·6 g/100 ml.

Discussion

We think that the unusual degree of interference with human folic acid metabolism found in this case was due to a combination of several factors. Firstly there was dietary deficiency of folic acid, secondly, extra utilization of folic acid due to marrow hyperplasia compensating for a haemolytic anaemia, thirdly, folic acid loss from the body by peritoneal dialysis, and finally there was the known antagonism of folate metabolism by the combination of trimethoprim and sulphamethoxazole.

The patient had been uraemic for 12 months and during this time ate poorly. This may well have critically depleted, if not have exhausted, her body stores of folic acid. She was then put on a Giovannetti diet containing 18 g of protein daily. Sevitt & Hoffbrand (1969) estimated that a 20 g protein diet supplies 150–300 μg of folic acid daily, after loss caused by cooking. They also demonstrated that peritoneal dialysis causes a significant loss of folate from the body.

As the estimated folic acid requirement is 50–100 μg daily (De Gruchy, 1970) it is obvious that the patient’s dietary folic acid intake was at a critical level. It is interesting to recall that Kahn (1968) showed in his study that haematological interference by co-trimoxazole could be prevented by 400 μg folic acid daily. Sharpstone (1969) showed that defective excretion of co-trimoxazole did not occur until creatinine clearance fell below 3·5 ml/min. Therefore, our patient whose clearance of creatinine was approximately 1 ml/min would have developed
drug retention at a time when her dietary intake was
below the level of 400 μg folic acid—the level shown
by Kahn to be necessary to counteract co-trimoxazole
antagonism of folic acid.

In addition to these factors, however, it may be
expected that her folic acid consumption was much
higher than normal as she had evidence of a signifi-
cant haemolytic anaemia. It is well known that a
haemolytic anaemia may cause megaloblastic
anaemia due to depletion of body folic acid stores—
Chanarin, Dacie & Mollin (1959). On the other
hand, Clarkson et al. (1970) did not comment on
megaloblastic anaemia in his cases with the haemoly-
tic uraemic syndrome.

Finally, during the later stages of her illness, she
had several peritoneal dialyses. Sevitt & Hoffbrand
(1969) showed that peritoneal dialyses caused a
striking fall in serum folic acid and this was un-
doubtedly a further significant factor in the develop-
ment of folic acid deficiency in our patient. One
must add that the low serum folic acid level does not
prove folate deficiency, as the bacteriological assay
may have been upset by the co-trimoxazole, although
it is strange that a similar mechanism did not
interfere with the estimation of vitamin B₁₂.

Conclusion

We would suggest that in this situation co-tri-
moxazole is administered with small doses of folic
acid and that, in fact, the Giovannetti diet itself might
be supplemented by a small daily supplement of
folic acid.

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G. M. Yuill

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