Correspondence

Role of ineffective erythropoiesis in anaemia of RA

Sir,

Samson et al. (1977) reported a study of the role of ineffective erythropoiesis in the anaemia of a patient with rheumatoid arthritis before and after gold therapy, suggesting that ineffective erythropoiesis may be an important reversible factor in the anaemia of rheumatoid arthritis. We feel that the results should be interpreted with caution.

The method of estimating ineffective erythropoiesis which these authors have used (Samson et al., 1976a) depends on a number of assumptions, the most important of which are (i) that the labelled compounds injected behave as tracers of the biological system. In the present study the amount of labelled δ-amino-α-ethylmalonic acid (ALA) administered was at least 1000-fold greater than the total body pool (Samson et al., 1976a) and cannot be considered a trace dose; (ii) that the mean normal ineffective erythropoiesis is 8%. This figure was taken from a study of plasma bilirubin kinetics (Berk et al., 1976) which in turn assumed that there was a normal degree of ineffective erythropoiesis in 20 patients with a wide variety of disorders. The constants derived from these assumptions in subjects with apparently normal erythropoiesis are further assumed to apply to patients with abnormal erythropoiesis; (iii) that the relative distribution of bilirubin derived from erythropoiesis and hepatic sources is the same in the plasma and in the faeces. However, plasma and hepatic bilirubin do not exchange completely and a significant proportion of hepatic bilirubin is excreted directly into the bile (Israel et al., 1963; Kirschenbaum et al., 1976).

It is perhaps not surprising that the results obtained using the method of Samson et al. (1976a) do not agree with estimates of ineffective erythropoiesis by other methods. Measurements of carbon monoxide production, bilirubin turnover, and iron kinetics have shown that mean normal ineffective erythropoiesis is approximately 20–30% (Coburn et al., 1963; Berk et al., 1970; Cook et al., 1970; Cavill et al., 1977). The method of Samson et al. (1976a), however, gave values between 4 and 12% in 4 normal subjects and gave a result which suggested that ineffective erythropoiesis in rheumatoid arthritis is three times the normal level. In 4 patients with classical or definite rheumatoid arthritis we have found, using a ferrokinetic method (Ricketts et al., 1975), that ineffective erythropoiesis is between 17 and 20% and does not significantly differ from normal.

In the present study (Samson et al., 1977) the data from one patient with rheumatoid arthritis described previously (Samson et al., 1976b) have been augmented by a partial study after gold therapy. The incorporation of labelled glycine and ALA into early labelled bilirubin was not measured. A change in ineffective erythropoiesis was inferred by assuming that hepatic haem turnover had remained constant after 2 years of therapy. No data were presented to justify this conclusion.

Measurement of the amount and effectiveness of erythropoiesis requires the use of sophisticated techniques. Under these circumstances it is easy to be beguiled by the results and their possible significance. However, the results are only as good as the methods by which they were obtained. We do not believe that the methods used by Samson et al. (1977) are based on physiologically and mathematically valid assumptions.

I. Cavill and C. Ricketts
Department of Haematology,
University Hospital of Wales,
Heath Park, Cardiff CF4 4XW.

This letter has been seen by Drs. Samson, Gumpel, and Halliday, who reply as follows:

Sir,

We have read the letter from Drs. Cavill and Ricketts with great interest. However, we feel that the complex points they raise regarding methodology would be more appropriately discussed in a haematological journal. The possible errors of our method and the criteria on which the results were concluded to be valid were stated briefly on our original paper (Samson et al., 1976a) and are fully discussed elsewhere (Samson, 1976). None of the comments made by Cavill and Ricketts has led us to modify our conclusions about either the method or this particular patient.

It seems most unlikely from morphological and cytochemical observations (Wickramasinghe, 1975), as well as from bilirubin and carbon monoxide data, that as much as 30% of normal erythroblasts die in the bone marrow. It is probable that even the 10%, or so of total bilirubin is derived from haemoglobin extracted with the nucleus rather than from cell death.

Contrary to the statement made by Cavill and Ricketts, the other authors to whom they refer do not support their high values for normal ineffective erythropoiesis. What Coburn et al. (1963) and Berk et al. (1970) showed was that 17–30% of haem turnover is normally derived from sources other than red cell destruction. This includes hepatic haem turnover as well as ineffective erythropoiesis and since the liver accounts for 13–20% of total haem turnover (Jones et al., 1971; Berk et al., 1976) only 4–10% can be derived from ineffective erythropoiesis. This would account for 5–12% of total erythropoiesis. Our results are thus in full agreement with these data.

The reason that the ferrokinetic method of Ricketts et al. (1975) gives much higher results may lie in their assumption that the late reflux of labelled iron arises entirely from cell death. There is no evidence that this is a valid assumption, and neither Pollycove and Mortimer
Penicillamine in seronegative polyarthritis

Sir,

In the letter entitled 'Failure of δ-penicillamine to affect peripheral joint involvement in ankylosing spondylitis or HLA B27 associated arthropathy' (Bird and Dixon, 1977) the authors report negative results in 7 patients with B27-associated arthritis who were negative for IgM rheumatoid factor. In my experience, 5 patients with classical ankylosing spondylitis treated with δ-penicillamine showed no therapeutic response, in complete accord with the reported findings. Further support for the lack of efficacy of penicillamine in ankylosing spondylitis has been published in France (Leca and Camus, 1975).

The authors then suggest that since there have been no placebo controlled trials which have shown that seronegative polyarthritis responds to penicillamine, it remains a possibility that seropositivity is a prerequisite for a favourable response. While it is correct that no such trial has been performed, many investigators engaged in the treatment of rheumatoid arthritis (RA) with penicillamine have reported characteristic favourable responses to penicillamine in seronegative patients who otherwise fulfill the criteria for RA. Albeit anecdotal, no one has reported that seronegative RA patients as a group are nonresponders. Although no placebo was used, penicillamine and gold were found to be equally effective in the management of seronegative juvenile chronic polyarthritis (Hall and Ansell, 1977). Surely every effort should be made in 'seronegative RA' patients to exclude B27 arthritis, systemic lupus erythematosus, sarcoidosis, bowel disease, etc. It would, however, seem unjustified to exclude a patient with classical RA from penicillamine therapy simply because of the absence of IgM rheumatoid factor.

ISRAELI A. JAFFE
1249 Fifth Avenue,
New York, NY 10029, USA

References

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I Cavill and C Ricketts

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