consider that the data support the presence of the disease in older children as well. Clearly, Bitot’s spots do not always indicate the presence of active xerophthalmia in children over 6, a point made in the original paper. The extent of severe malnutrition in all age groups prompted me to examine children up to 14 in this population. It was an unprecedented situation, which I thought merited a wider sample size than normally used.

While Dr McLaren thinks that corneal xerosis cannot be diagnosed without a slit lamp, numerous other investigators have reported corneal xerosis in prevalence surveys performed without a slit lamp. In fact, the WHO ‘Field guide to the detection and control of xerophthalmia’ states that ‘(corneal xerosis) is easily diagnosed and highly specific’ and makes no mention of the use of a slit lamp to make this diagnosis.1 I feel confident in the diagnosis as stated in the paper.

Finally, and most disturbing, is the point made by Dr McLaren that while the WHO guidelines are not engraved on stone tablets, they should stand until new data become available. The point of this paper is to suggest that in severe disaster situations just such a revision may be indicated. It is time for this issue to be discussed in an open forum to prepare for the next disaster, which most unwelcomely and assuredly will arise.

Reference

Meningitis presenting as hypertension

Sir,

I have just read the report by Waters and Gillis of meningitis presenting as hypertension.1 A month ago I was called urgently to see a child with haemophilus meningitis who had developed severe hypertension. The hypertension was cured instantly by using an appropriately sized sphygmomanometer cuff. Reports involving blood pressure measurement in children should always give precise details of how the blood pressure was measured.

Reference

D P Addy
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Meningitis due to Haemophilus influenzae type b resistant to ampicillin and chloramphenicol

Sir,

The recent report by Fraise et al of multiply resistant Haemophilus influenzae1 makes reference to two previous reports of resistance found in this organism from the United Kingdom2 and the United States.3 I suspect the authors did not take the time to read the reference for the USA. If they had it would have been clear that it was describing resistance in Australia, a country not easily confused with the USA. Even correct reading of the title should have made the error apparent.

Chloramphenicol resistance is now widespread, including reports from Thailand,4 the USA,5 and the Caribbean.6

Readers are entitled not only to accuracy of data but also to diligence and care of presentation.

References

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Drs Fraise, Mecks, and Richards comment:
We have read with interest the comments made by Dr Forsyth. We acknowledge that there was an omission in the introductory phrase of our paper, which should have read ‘... in the United States and in Australia.’ We do not believe a small omission such as this, however, changes in any way the validity of our report or the message it contains.

This correspondence is now closed—ed.

Thrombotest values and effect of vitamin K administration for infants

Sir,

The finding of Garrow et al that Thrombotest values decline on the second day of life in breast fed babies but not in those who have received vitamin K at birth is important.1 Fetal haemorrhagic episodes, including intracranial haemorrhage, are more common at 2 to 3 weeks of age than in the early days of the neonatal period.2 Thus the Thrombotest for babies at 2 weeks may also be useful to predict and manage vitamin K deficiency.

We performed the Thrombotest (Thrombotest Owren, Eisai Co, Tokyo) for 437 babies at 2 weeks. Of the 437 babies, 263 were breast fed, 37 bottle fed, and 137 mixed

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