retention of urine with overflow, faecal incontinence, drowsiness, and neck stiffness. 17 days from the onset of loss of vibration sense, diminution in joint position sensation, and a qualitative loss of sensation to pinprick had developed, involving both legs and extending to the area supplied by the 8th cervical spinal segment. Some return of power in the legs was noted on day 19 of illness, and over the next week there was a steady improvement. 6 weeks later she had only slight weakness of the legs, but after 6 months there were no abnormal signs, and her school reported her academic ability apparently unimpaired.

Lumbar puncture after admission produced clear cerebrospinal fluid under normal pressure containing 15/mm³ white cells, mainly mononuclear, protein 0·36 g/l, and glucose 2·5 mmol/l (45 mg/100 ml), which was sterile on culture. A further tap a few days later showed a slightly higher white cell count (46/mm³, 90% lymphocytes) but a lower protein. A marked neutrophil leucocytosis (total white cell count 41·0 × 10⁹/l (41000/mm³) with 87% polymorphs) was present on admission and persisted for 4 weeks. A thorough and repeated search for bacterial infection including a bone scan was negative. Acute and convalescent antibody titres done in parallel did not suggest infection with Brucella, Epstein-Barr virus, Rickettsia, adenovirus, mumps, cytomegalovirus, measles, herpesvirus, Toxoplasma, Leptospira, Staphylococcus, or Salmonella typhi and paratyphi. Coxsackie antibody titres also were normal except for B₃ which showed a raised titre on admission (1:1024), falling to 1:256 by day 19 of illness. Viral cultures were negative.

We feel this very high Coxsackie B₃ titre suggests that the illness was caused by this organism, and we cannot find any published reports of this combination, though transverse myelitis has been described in association with B₃ virus (Dery et al., 1974). The persistent neutrophil leucocytosis, which was a worrying feature of the illness, does not seem to have been a feature of the rare cases of transverse myelitis with other viral aetiology.

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Reference

Maternal diabetes and congenital malformation

Sir,

May I add to the experience of Drs. Day and Insley detailed in their paper on this subject (Archives, 1976, 51, 935), as it is very similar to my own (Komrower, 1961). I followed 213 consecutive pregnancies where all the mothers had received insulin therapy. There were 20 deaths before the 28th week (14 less than 20 weeks), 16 stillbirths and 20 neonatal deaths. I was able to review 182 children on more than one occasion, some for more than 10 years, and by good chance I adopted similar definitions to Day and Insley for major and minor malformations.

There were 19 significant congenital abnormalities: 5 in stillbirths, 3 in neonatal deaths, and 11 in living children. The system involvement was similar to that described by the Birmingham authors—cardiovascular 5, skeletal 5, urogenital 4, oesophageal atresia 1, central nervous system 2 (one child had both anencephaly and multiple skeletal deformities), and 3 had large haemangiomas. The incidence was 9·75% compared with a hospital figure of 3·2% for a 2-year period during the study (the latter results were obtained from our monthly neonatal report). Unfortunately, I do not have the ages of the mothers concerned but the duration of their diabetes varied between 6 months and 10 years. This study took place before our present experience in the treatment of hyaline membrane disease but even so one was able to show that a diabetic woman reaching the 28th week of her pregnancy had a 75% chance of producing a healthy baby without a congenital abnormality.

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Reference

Fibrosing alveolitis

Sir,

Dr. Hewitt and his colleagues, in their paper on fibrosing alveolitis in infancy and childhood (Archives, 1977, 52, 22), state that all but 2 of their 10 children improved when given corticosteroids, and in their summary they further state that in children fibrosing alveolitis is almost always a corticosteroid-responsive disease. This being the case, I should be interested to learn why 5 of their patients died of the disease, including 4 shown in Table 8 to be steroid-responsive.

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Prof. D. Hull and Dr. C. J. Hewitt comment:

We anticipated that some readers of the article might raise questions that could be more appropriately answered by full detailed case histories. We would be prepared to send these direct to any interested clinician.