Late onset ornithine carbamoyl transferase deficiency in males

Sir,

We were interested that Drogari and Leonard did not mention sodium valproate as a possible precipitant of symptoms in late onset carbamoyl transferase deficiency in males.1 We wish to report such a case; to our knowledge this has not been reported before.

A 10 year old boy, who had no previous medical or family history, was started on sodium valproate 200 mg twice a day for worsening photosensitive absence seizures. After three weeks a subtherapeutic concentration of the drug prompted an increase in dose to 200 mg three times a day. Five days later the boy vomited twice. Solids and sodium valproate were withheld. The next day he vomited three more times and became difficult to rouse in the general practitioner's surgery. He was immediately admitted to hospital but remained easily rousable overnight. Twelve hours after admission his conscious level deteriorated. Blood glucose concentration was 5.5 mmol/l and ammonia 145 μmol/l, his prothrombin ratio was 1.5 and aspartate aminotransferase activity 47 IU/l. All other liver function tests and a computed tomogram were normal. Blood sodium valproate concentration was 0.16 mg/l (subtherapeutic).

Eighteen hours after admission he was transferred to another hospital for intensive care. Assisted ventilation was begun. Intracranial pressure was low initially. Thirty six hours after his first hospital admission the blood ammonia concentration was 245 μmol/l and intracranial pressure 2.67–4.00 kPa. Chromatography of the urine showed a peak of orotic acid. This was initially a very small peak but after reversal of the effect of sodium valproate with sodium benzoate, became a very large peak typical of ornithine carbamoyl transferase deficiency. Despite aggressive management of his intracranial pressure and appropriate doses of dextrose, sodium benzoate, arginine, and growth hormone, intracranial pressure rose to 7.33 kPa and ammonia to 918 μmol/l over the next 12 hours before coming under control. His cerebral perfusion pressure could not subsequently be maintained and he died on the sixth hospital day. Very low residual ornithine carbamoyl transferase activity of 1·6 μmol/mg/hour (normal carbamoyl phosphate synthetase activity) was confirmed in liver obtained immediately postmortem.

Hyperammonaemia and brain swelling in valproate toxicity complicating urea cycle disorders are more often rapidly progressive and fatal than in classical Reye's syndrome even if, as in the present case, early and aggressive treatment is given. Such deaths are one of several metabolic hazards of sodium valproate that must be weighed against the potential benefit of treatment.2

References


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Rubella immunisation for girls over 14 years

Sir,

I was interested to read Dr Hodes' article on rubella immunisation for girls over 14 years,1 but believe that her findings may not justify her recommendation that rubella vaccination should be offered to girls until they leave school at 16 years of age.

Dr Hodes offered her programme to 56 of 83 girls whose rubella immunity status was unknown, but she does not tell us why the other 27 were not invited. Only 25 of the 56 accepted, and 21 of them (84%) were found to be seropositive. The other four were vaccinated. The increase in the proportion of girls vaccinated or immune from 86% to 90% was due largely to the finding of seropositive girls, and much less to vaccination.

If it is assumed that the same proportion (84%) of all 83 girls was immune, 566 girls (96-6%) were vaccinated or immune before Dr Hodes' programme was introduced, and 570 (97-2%) afterwards. While applauding every effort to increase rubella immunity in the community, I wonder whether this marginal increase justified the expense of the enterprise.

Reference


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Dr Hodes comments:

Professor Smithells rightly criticises the fact that only 56 of the 83 eligible girls were invited. This was due to many practical problems that were described in the second and third paragraphs of the discussion. This is a shortcoming of the study and in future one would have to try harder to invite all eligible girls.

Assuming that the same proportion (84%) of all 83 girls was immune, 70 girls would be seropositive and 13