shown a shortage of postnatally dividing microneuromes in the cerebellar granular layer and other neurones in the deeper layers of the cerebral cortex. The deficit is particularly surprising in view of the comparatively mild growth retardation imposed particularly surprising in view of the comparatively mild growth retardation imposed during the brain's vulnerable period of development. The comparable period of human brain growth extends from about 30 weeks of gestation into the second year of postnatal life. Behavioural correlates in the experimental animal were also discussed.

From mouse to man? R. D. Barnes (Division of Infant Development, Clinical Research Centre, Northwick Park, Harrow). The techniques of ovum transplantation and ovum fusion are described since these have been successfully used to investigate various diseases in mice. Ovum transplantation allows the transfer of one fertilized egg to an animal of another strain and is a very useful technique to investigate the role of transplacental infection in any disease. Secondly, ovum fusion derived chimaeras provide an excellent experimental model to investigate the interrelation between two cell populations in the same animal. Furthermore, in this respect ovum fusion allows us to investigate the hypothesis that certain diseases in mice, including autoimmune haemolytic anaemia and leukaemia, occur due to the time-related failure of what might be considered a normal recognition and inhibition control process, and evidence for this is presented.

Finally, the modification and further sophistication of in utero grafting techniques might, in the event of proving useful in curing certain defined diseases in mice, suggest that these techniques could subsequently apply to man. However, this must await full investigation in animals, and furthermore, the development of suitable techniques for antenatal diagnosis of affected children.

Coagulopathy in the hypoxic newborn baby. M. A. Chadd (introduced by O. P. Gray) (Department of Child Health, Welsh National School of Medicine). It is well established that coagulation defects and haemorrhage may result in death in the perinatal period.

Classical haemorrhagic disease as first described in the 1800s by Townsend is now of almost academic interest with the decline in breast feeding and the reduced use of vitamin K in neonatal nurseries.

The concept of secondary haemorrhagic disease first proposed by Aballi and de Lamerens 10 years ago is recognized as becoming of increasing importance. It is frequently found in association with hypoxia and is unresponsive to vitamin K.

This study of 75 hypoxic newborn and 75 control infants was undertaken in an attempt to elucidate the nature and incidence of coagulation defects in the newborn and the role of hypoxia in producing such defects.

Brain oedema induced by asphyxia in newborn rats. S. W. D'Souza (introduced by J. A. Davey) (Department of Child Health, University of Manchester). Using 5-day-old rats an attempt has been made to induce brain oedema experimentally with asphyxia as the precipitating insult.

Changes in brain water, sodium, and potassium content can be induced by slow prolonged asphyxia at body temperature and to a lesser extent in the cold, but not by acute anoxia. These changes are more marked in the brainstem than in the hemispheres.

Plasma growth hormone response to intravenous glucagon administration. L. Stimmilger and G. Snodgrass (Guy's Hospital, London). Glucagon administration has been shown to stimulate growth hormone secretion. In a previous study sampling times were delayed, whereas it is known that glucagon produces a very rapid rise in plasma insulin.

Glucagon 15 μg/kg body weight was administered intravenously to a group of 24 children being investigated for abnormally short stature. Blood samples were obtained immediately before and at frequent intervals after glucagon administration. 20 of these children showed a marked rise in growth hormone levels. Of these patients a significant rise had already occurred at 2 minutes after glucagon administration. The time of peak growth hormone response occurred before 20 minutes in 17 of these patients. Insulin and glucose levels were also estimated. The interrelation between these parameters and the growth hormone response was discussed.

Insulin release from human fetal pancreas in vitro. R. D. G. Milner and M. A. Ashworth (Department of Child Health, University of Manchester). Pieces of pancreas removed from dead human fetuses delivered by hysterotomy were incubated in vitro as described previously for rabbit pancreas. The gestational age of the fetuses studied was between 14 and 24 weeks and their body weights ranged from 50 to 625 g. Insulin released into the incubation medium was measured under basal conditions and in the presence of various substances known to stimulate insulin secretion in adult man and other species. Glucose (3-0 mg/ml) did not stimulate insulin release. Stimuli which are thought to act by raising intracellular levels of cyclic AMP: glucagon (5 μg/ml), theophylline (1 mM), and dibutyryl cyclic AMP (1 mM) stimulated insulin release in the presence of 0.6 or 3.0 mg/ml glucose or in its absence. Ionic stimuli which act late in the stimulus-secretion pathway were also uniformly effective: barium (2.54 mM), ouabain (10⁻⁵ M) and potassium (60 mM). Leucine (5 mM) and arginine (5 mM) were effective in some experiments only. Tolbutamide (400 μg/ml) was ineffective in two experiments.

It was concluded that cells capable of secreting insulin are present in human fetal pancreas from the 14th week of fetal life onwards.

Applications of praeordial accelerometry. D. Pickering (introduced by B. D. Boxer) (Department of Paediatrics, Radcliffe Infirmary, Oxford). A praeordial accelerometer is described which has been
constructed from silicon semiconductor strain gauge, which has a linear response from DC to 1000 c.p.s. Its applications are described in relation to phonocardiography, to the timing of cardiovascular events, to cardiac output, and lastly as a guide to the force of myocardial contraction. The small size of pick-up and high frequency response make it a useful tool for phonocardiography in childhood and the applications to the recording of systolic clicks associated with small ventricular septal defects are described. The use of an accelerometer to calibrate the time lag of pressure waves up a Lehmann catheter is described and the correlations of the acceleration waves with intracardiac pressures are shown. The relation of ejection time to cardiac output in congenital heart disease is discussed and finally evidence is shown that the amplitude of the accelerogram waves is related to the force of myocardial contraction. A pendulum method of calibration and further applications in the monitoring of myocardial disease are suggested.

Experience with treatment of covert bacteriuria in 5-year-old Dundee schoolgirls. D. C. L. Savage, M. E. Wilson, M. McHardy, and W. M. Fee (Departments of Child Health and Bacteriology, University of Dundee, and Child Health Services, City of Dundee).

Since 1967 the 5-year-old schoolgirl entrants to Dundee Primary Schools have had their urine examined for significant bacteriuria. Approximately 5000 children have been screened and a prevalence of 1·5% covert bacteriuria has been found.

Forty children detected in the first 2 years screening were all treated and the results of treatment, follow-up, and repeat radiological investigation 2 years later were presented. The most significant finding was the high rate of reinfection while on therapy, which was not influenced by the presence or absence of underlying urogenital abnormality. At 2 years over 75% had become reinfected on at least one occasion and over half the children had had 2 or more episodes of re-infection. It was unusual to find an acute illness associated with these episodes, though in many cases mild urinary symptoms recurred. In only one child does the radiological picture and renal function give cause for grave concern.

Children detected more recently, approximately 40, have entered a randomized controlled trial of therapy. The follow-up period is still brief but a number have been observed for over a year. It is already apparent that some resolve without therapy and in no child has serious symptomatic disease developed.

Detection of heterozygotes for homocystinuria by oral loading with L-methionine. I. B. Sardharwalla, B. Fowler, and A. J. Robins (introduced by J. B. Houston) (Royal Manchester Children's Hospital and Department of Medical Biochemistry, University of Manchester). To be published elsewhere.

Some consequences of artificial feeding in neonates with reference to excess weight gain and osmolar loading. L. S. Taizt (introduced by V. Dubowitz) (Department of Child Health, University of Sheffield). Recent studies in Sheffield have indicated that infants showing a rapid rate of weight gain in early infancy have a greater tendency to later obesity than those who gain weight more slowly.

These findings have prompted further studies of feeding practices. It has been found that rapid weight gain is associated with artificial feeding and the early introduction of solids. On the basis of Eid's criteria 59% of infants show excessive weight gain at 6 weeks. These findings are associated with an estimated dietary intake that exceeds the usually recommended 100 calories/kg per day.

Analysis of milk samples taken from bottles brought to the follow-up clinic show that the sodium concentration often exceeds that of cow's milk, indicating that insufficient care is taken in the preparation of feeds.

This increased osmolar intake may be significant in relation to the high incidence of hypertonic dehydration.

Clinical value of plasma creatine kinase and uric acid levels during first week of life. B. A. Wharton Urmilla Bassi, G. Gough, and Angela Dilhams (Queen Elizabeth Hospital, London E2). Published in full (Archives of Disease in Childhood, 46, 356).

Demonstrations

Anonymous mycobacteria in childhood. T. Knowlson, W. A. Hyde, and H. B. Marsden (Royal Manchester Children's Hospital).

Tumours in children. J. K. Steward (Manchester Children's Tumour Registry).

Some problems posed by the sweat test. V. Schwarz (Department of Child Health, University of Manchester).

Studies on the mechanism of sweat secretion. C. Gordon and V. Schwarz (Department of Child Health, University of Manchester).

Vulnerability of the developing brain. (Department of Child Health, University of Manchester): (a) Effects of experimental growth retardation—Jean Sands and J. Dobbing; (b) Brain enzymes following experimental undernutrition—B. Adlard; (c) Behavioural consequences: of experimental undernutrition—J. Smart and A. Lynch; (d) Experimental X-irradiation in infancy: effects on brain enzymes and behaviour—B. Adlard and A. Lynch; (e) Human brain growth—Jean Sands and J. Dobbing.
Applications of praecordial accelerometry.

D Pickering

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