Douglas Gairdner won a scholarship to Oxford before transferring to medicine at the Middlesex Hospital, London, graduating in 1936. After a fellowship at Bellevue Hospital, New York, he served with the Royal Army Medical Corps in the Middle East. After the war he was appointed assistant to Professor Sir James Spence in Newcastle before moving to Cambridge in 1947. His contributions to paediatric research and literature, which his modesty prevents him from enumerating, are too extensive to detail. His approach is exemplified, however, when he says of neonatal research 'It was a joy for us that the extremely simple technology, which was all we had, was yet capable of yielding worthwhile results' and he praises the dedication of his hard working colleagues. He was one of the many paediatricians of the day who rejoiced in the elevation of babies from being something of second class citizens to fully accepted 'patients' in their own right.

Douglas Gairdner has served all aspects of paediatrics with great distinction and is possibly best known for his 15 years Co-Editorship of the Archives of Disease in Childhood and Editorialship of the first four editions of Respiratory Advances in Paediatrics. The BPA recognised his many contributions in awarding him the Spence Medal (1976) and electing him to Honorary Membership (1977).

C H M WALKER

LETTERS TO THE EDITOR

Mechanical ventilation and respiratory syncytial virus infection

Sir,—Drs Lebel et al report weight and prematurity as risk indicators for the need for ventilation in bronchiolitis.1 In their case-control study a viral cause was determined in only 32%; infants with underlying cardiopulmonary abnormalities—who are known to be at increased risk for severe respiratory syncytial virus infections and mortality2—were excluded. Thus the results from their study may not readily be applied to all infants with bronchiolitis who need hospitalisation.

We would like to present the preliminary results of a prospective study (1987-9) of children admitted with a respiratory syncytial virus infection to the Sophia Children’s Hospital, The Netherlands. In all patients the infection was proved by direct immunofluorescent assay. Nineteen patients needed mechanical ventilation and 82 were admitted to the general ward.

Weight and age (corrected for prematurity) on admission were significantly related with mechanical ventilation (by χ², p<0.05). No relation between mechanical ventilation and risk factors for severe respiratory syncytial virus infections (prematurity, congenital heart disease, bronchopulmonary dysplasia, or immunodeficiency states) was found. Nine (47%) of the ventilated patients and 36 (43%) of the non-ventilated patients belonged to the risk group. In a stepwise logistic regression analysis (while controlling for the covariates: age, prematurity, and risk factors for severe respiratory syncytial virus infection) only weight appeared to be significantly related to the need for mechanical ventilation (coefficient −0.36682 (SE 0.1685), p=0.01), while none of the other variables did. Prematurity and age were not independent risk indicators for mechanical ventilation in this study. In contrast to Lebel et al, who report odds ratios as ‘relative risk’, from the data of our prospective study we are able to estimate the absolute risk of mechanical ventilation related to the weight on admission (figure). A low weight is related to an increased risk for mechanical ventilation. For infants weighing less than 5000 g, a relative risk for mechanical ventilation of 4.3 (95% confidence interval 1.3 to 13.9) was estimated.

H A VAN STEENKEL-MOLL W J E TISSING

Department of Pediatrics, Sophia Children’s Hospital, Erasmus University Rotterdam, Gerebrugel 160, 3038 GE Rotterdam, The Netherlands

J A HAZELZET

Department of Pediatrics, Subdivision of Puerperal Intensive Care, Sophia Children’s Hospital, Erasmus University Rotterdam, Gerebrugel 160, 3038 GE Rotterdam, The Netherlands


3 MacDonald NE, Brees Hall C, Sufin SC, Alexson C, Harris PJ, Manning J. Respiratory syncytial virus infection in infants.

Helicobacter pylori and protein losing enteropathy

Sir,—We have demonstrated a high prevalence of anti-Helicobacter pylori (formerly Campylobacter pylori) antibodies in children under the age of 3 years in The Gambia, West Africa.1 Using a serological test validated by histology and microbiology, 47/77 (53%) of children with chronic diarrhoea were shown to have significantly high anti-H pylori IgG antibody titres, and indeed, the prevalence of H pylori antibodies in healthy asymptomatic children of the same age was also high at 26%.

It was against the background of these findings that we were interested in the description, in 1987, by Hill and colleagues, of transient protein losing enteropathy in association with acute infection with H pylori.2 This observation might be due to chance but could have important consequences in the nutritional rehabilitation of infants with chronic diarrhoea and severe protein energy malnutrition. Therefore we undertook a study to establish whether or not H pylori infection was associated with protein losing enteropathy in Gambian children with chronic diarrhoea.

Fifty three subjects (25 boys, 28 girls; mean age 19 months) were studied and all had chronic diarrhoea (more than three stools per day for more than two weeks) and severe protein energy malnutrition (32 marasmus, 21 marasmic-kwashiorkor). After admission, three consecutive fresh stool samples were collected for serological, parasitological, and bacteriological investigation. Specific anti-H pylori IgG antibody was measured by enzyme linked immunosorbent assay (ELISA) in all subjects. Gastroscopy and antral mucosal biopsy were performed in 20/53 children.

Protein losing enteropathy was estimated by random faecal α1-antitrypsin measurement. Whole single stools were collected ensuring that both the liquid and solid phase were obtained. These were frozen and lyophilised and α1-antitrypsin measured by a single radial immunodiffusion method. Random faecal α1-antitrypsin measurement (normal mean (SD) value in healthy Gambian children 1.54 (0.23) mg/g stool) has been shown to be a reproducible screening test for excessive enteric protein loss and has been validated against 31 chromium labelled albumin excretion in the stool.3

Fifty six percent of this group of children had significantly raised anti-H pylori antibody titres and in 11/20 gastroscopically this was associated with recovery of the organism and histological gastritis. Strongyloides stercoralis, and Giardia lamblia were found in 11 and 38% of the patients respectively. Hypoaflagoria occurring in children with S stercoralis was found to be associated (r=0.952, P<0.001) with increased faecal α1-antitrypsin excretion (mean (SEM)=2.47 (0.9) mg/g stool) whereas there was no difference between levels of faecal α1-antitrypsin found in children with 1.57 (0.2) mg/g stool or without 1.62 (0.3) mg/g stool evidence of H pylori infection.

Therefore, in this study we failed to show that chronic infection with H pylori is associated with protein losing enteropathy.

P B SULLIVAN

Department of Child Health, Westminster Children’s Hospital, London SW1P 2NS

J E THOMAS E J EASTHAM

Department of Child Health, University of Newcastle upon Tyne

P G LUNN G NEALE

Dun’s Nutritional Laboratory, Cambridge

Mechanical ventilation and respiratory syncytial virus infection.

H A van Steensel-Moll, W J Tissing, M Offringa and J A Hazelzet

Arch Dis Child 1990 65: 332
doi: 10.1136/adc.65.3.332

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