Early report

lleal-lymphoid-nodular hyperplasia, non-specific colitis, and pervasive developmental disorder in children

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Summary

Background We investigated a consecutive series of children with chronic enterocolitis and regressive developmental disorder.

Methods 12 children (mean age 6 years [range 3-10], 11 boys) were referred to a paediatric gastroenterology unit with a history of normal development followed by loss of acquired skills, including language, together with diarrhoea abdominal and pain. Children underwent gastroenterological, neurological, and developmental assessment and review of developmental records. Ileocolonoscopy and biopsy sampling, magnetic-resonance imaging (MRI), electroencephalography (EEG), and lumbar puncture were done under sedation. Barium follow-through radiography was done where possible. Biochemical, haematological, and immunological profiles examined.

Findings Onset of behavioural symptoms was associate by the parents, with measles, mumps, and rub vaccination in eight of the 12 children, with measl infection in one child, and otitis media in an All 12 children had intestinal abnormalities from lymphoid nodular hyperplasia to a ration. Histology showed patchy chronic inflan. perplasia in in 11 children and reactive ilea mpho s included seven, but no granulomas. Be vioural disor autism (nine), disintegrative sychologis (one), and possible postviral or vaccinal encephalitis (o). There were no focal neurological ab malities and and EEG tests were normal. Abnor al laboratory results are significantly thylmal compared with ageraised urinary 03), low haemoglobin in four matched control m IgA in ar children. children, low s

Interrectation be idented associated gastrointestinal discusse and revelopmental regression in a group of previously amarchism, which was generally associated in time to possible environmental triggers.

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Introduction

We saw several children who, after a period of apparent normality, lost acquired skills, including communication. They all had gastrointestinal imptoms, including abdominal pain, diarrhoea, and mating and, it some cases, food intolerance. We discribe the clinical follings, and gastrointestinal feature of these charges.

Patients and metitals

12 children, cons ativel red to a his paediatric gastre y of a pervasive rerology developmental der with loss ed skills and intestinal abdominal ain, bloating and food symptoms arrh gated. All children were admitted to the intolerance), were inve ward for week, accomp ed by their parents.

(Inical investigations

took historic including details of immunisations and eleasure to infect as diseases, and assessed the children. In 11 case the history as obtained by the senior clinician (JW-S). Neuroccially and psychiatric assessments were done by sonsultant staff (PH, MB) with HMS-4 criteria. Developmental records from purents, health visitors, and general practitioners. Four children did not undergo psychiatric assessment in hospital; all had been assessed professionally elsewhere, so these assessments were used as the basis for their behavioural diagnosis.

After bowel preparation, ileocolonoscopy was performed by SHM or MAT under sedation with midazolam and pethidine. Paired frozen and formalin-fixed mucosal biopsy samples were taken from the terminal ileum; ascending, transverse, descending, and sigmoid colons, and from the rectum. The procedure was recorded by video or still images, and were compared with images of the previous seven consecutive paediatric colonoscopies (four normal colonoscopies and three on children with ulcerative colitis), in which the physician reported normal appearances in the terminal ileum. Barium follow-through radiography was possible in some cases.

Also under sedation, cerebral magnetic-resonance imaging (MRI), electroencephalography (EEG) including visual, brain stem auditory, and sensory evoked potentials (where compliance made these possible), and lumbar puncture were done.

Laboratory investigations

Thyroid function, serum long-chain fatty acids, and cerebrospinal-fluid lactate were measured to exclude known causes of childhood neurodegenerative disease. Urinary methylmalonic acid was measured in random urine samples from eight of the 12 children and 14 age-matched and sex-matched normal controls, by a modification of a technique described previously. Chromatograms were scanned digitally on computer, to analyse the methylmalonic-acid zones from cases and controls. Urinary methylmalonic-acid concentrations in patients and controls were compared by a two-sample t test. Urinary creatinine was estimated by routine spectrophotometric assay.

Children were screened for antiendomyseal antibodies and boys were screened for fragile-X if this had not been done