Interestingly, serum creatine kinase BB measured by radioimmunoassay, was abnormal in this patient (figure). Brain-type creatine kinase BB is present at a lower concentration, in other tissues. In cases of proved macroscopic brain injury very high enzyme levels may be detected within hours. Here, highest levels were observed after 48 hours, possibly when hypoxic cells developed structural changes due to continuing ischaemia. Previously we found transient abnormalities of creatine kinase BB and MB in marathon runners without neurological deficits. In this instance creatine kinase MB was normal while the BB isoenzyme remained high and then fell, correlating with clinical improvement.

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Outpatient paediatric fibreoptic proctosigmoidoscopy: possible and useful

The role of colonoscopy in diagnosis and follow-up of children with chronic inflammatory bowel disease is becoming recognised. In adult practice flexible fibreoptic sigmoidoscopy without sedation and with minimal bowel preparation is a safe and useful investigation. In assessing adult inflammatory bowel disease conventional rigid proctosigmoidoscopy and rectal biopsy are normally the first-line procedures. In most paediatric centres, however, rigid proctosigmoidoscopy and rectal biopsy are performed only under general anaesthesia and are regarded as major investigations, rarely undertaken. We therefore decided to see if limited fibreoptic proctosigmoidoscopy using a small-diameter colonoscope would be acceptable to children in a paediatric outpatient clinic and to evaluate its role in the diagnosis and follow-up of children with inflammatory bowel disease.

Patients, methods, and results

Fibreoptic proctosigmoidoscopy was attempted in 21 children (13 boys and 8 girls, age range 4-15 years, mean age 11) attending the paediatric inflammatory bowel disease clinic at St Bartholomew's Hospital between February and May 1982. Fibreoptic proctosigmoidoscopy was added without formality to the routine outpatient physical examination. No prior bowel preparation was given and no sedation was used before, during, or after the procedure. A 1 cm diameter very flexible paediatric colonoscope with tip designed for comfortable insertion (Olympus PCF) was selected. Patients were examined in the left lateral position. Only limited examination was attempted. Biopsy samples for histological assessment were normally taken at 10 cm intervals, at least one biopsy specimen being taken from the rectum during each examination.

The instrument was successfully inserted to between 20 and 30 cm in 20 of the 21 patients. The time for the whole procedure including taking biopsy specimens and teaching was 2-10 minutes (mean 5-6 minutes). One examination was impossible because of impaction of solid faeces. Fifteen of the children felt no discomfort and the remaining five tolerated the procedure, though finding it initially uncomfortable.

In four patients the tissue samples taken were considered inadequate for histological assessment, mainly because of their small size. Nine children showed histological abnormalities, including three in whom the mucosa appeared normal endoscopically. In six patients the results of the examinations were thought to have aided clinical management. Five of the patients subsequently underwent total colonoscopy with bowel preparation, in each case giving results compatible with those obtained during the limited examination.

Comment

This small study shows that fibreoptic proctosigmoidoscopy can be quickly performed in a routine paediatric outpatient clinic and is acceptable to children. It allows the paediatric gastroenterologist to follow the standard practice in adult clinics of rapid inspection of the rectal mucosa and the taking of a rectal biopsy specimen. The paediatric colonoscope is smaller in diameter than an examining finger and is considerably more comfortable than the rigid metal instrument. The colonoscope and its light source can be conveniently mounted on a standard metal dressing trolley (figure). The "teaching examination" side arm allowed some children and parents to appreciate the nature of the clinical problem by watching during the examination, as well as being convenient for discussion with students or medical colleagues. Photographic documentation was easy for those who wished to take the image to the bedside as a teaching aid.

The only drawback of the paediatric colonoscope used for this study was the small-size forceps biopsy specimens obtained. The importance of taking biopsy specimens is considerable; thus multiple samples should be taken in each case.

Though fibreoptic proctosigmoidoscopy gave useful information in these cases, it cannot supplant the additional information given in some cases by total colonoscopy. The occurrence of rectal bleeding in Crohn's disease means that a very limited examination with normal biopsy appearances might be misleading, though in our experience there is frequently minor abnormality in the sigmoid colon which is visible on limited examination. Small size is offset by the taking of accurate "target" specimens, which give a high percentage of successful histological diagnosis of Crohn's disease. Whereas total colonoscopy requires considerable experience which may not be available in every paediatric centre, limited examination is very easy to perform and requires little training. Our study convinced us that paediatric fibreoptic proctosigmoidoscopy without sedation or bowel preparation is practicable, well tolerated, and a useful investigation in diagnosis and management of children with possible inflammatory bowel disease.

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