Computed tomography in chronic inflammatory bowel disease

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Abstract

In children with complicated inflammatory bowel disease, conventional ultrasound imaging may not define the extent of extraluminal disease and the involvement of other viscera. Three children with chronic inflammatory bowel disease are presented, where computed tomography was well tolerated and provided valuable information on extraluminal disease, involvement of other organs, and the state of the bowel wall and mesentery. In children in whom ultrasound examination is inconclusive or limited by gas or tenderness, computed tomography can provide important information that may determine clinical management.

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Initial diagnosis of inflammatory bowel disease in children requires a combination of barium studies, endoscopy, and biopsy to define the extent of disease. In severe ulceration, transmural disease may result in mesenteric changes and extraintestinal complications which cannot be defined by barium studies. Cross sectional imaging is required to define the extent of extraluminal disease to determine management. In the paediatric patient, ultrasound is the first line investigation, but not infrequently fails to define the extent of the clinical problem fully. In these circumstances, computed tomography has been shown in adult patients to influence management; however, there are only limited reports of its use in paediatric practice. We present three children with inflammatory bowel disease in whom computed tomography contributed valuable information and influenced clinical management.

Case reports

CASE 1
A 16 year old boy was admitted with an acute history of abdominal pain, fever, and anorexia. He had a nine year history of ileocaecal Crohn’s disease and had been treated with a right hemicolectomy at the age of 12 years. He subsequently developed an anastomotic relapse which required further small bowel resection. Before this episode, he had been in remission for two years.

On admission he was tachycardic and dehydrated. Investigations showed a haemoglobin of 88 g/l, white blood count (WBC) $24 \times 10^9/l$, platelets $638 \times 10^9/l$, erythrocyte sedimentation rate (ESR) 69 mm/h, albumin 29 g/l, and C reactive protein 115 mg/l. Plain abdominal x ray showed moderate proximal constipation, but no small bowel dilatation. Ultrasound examination revealed a thick walled rectum but no evidence of a fluid collection. Sigmoidoscopy under anaesthesia showed moderate rectal inflammation. He was treated with intravenous antibiotics, corticosteroids, and total parenteral nutrition. Despite this, he remained very unwell and developed a swinging pyrexia to 39°C.

Because of concern about the presence of an occult abscess, computed tomography was performed to assess the rectum and perineum further. This showed gross thickening of the sigmoid colon and rectal wall with complete loss of the perirectal fat planes, and adherence of the rectum to the bladder. Despite the extensive inflammatory change no discrete pelvic collection was seen (fig 1A and B). A diversion colostomy was performed to defunction the area of diseased bowel, with the rectum left...

Figure 1 Case 1. Computed tomography of pelvis with rectal contrast, showing (A) marked inflammation of perirectal tissue with loops of small bowel and sigmoid colon adhering to the rectum (arrow), and (B) gross asymmetrical thickening of the rectal wall and posterior wall of the bladder (arrows).
in situ. The postoperative period was complicated by a rectovesical fistula which settled spontaneously.

Two months after surgery, repeat computed tomography showed dramatic improvement, with restoration of the pelvic fat planes, reduction in bowel wall thickening, and no evidence of rectovesical fistula (fig 2A and B). He remains in remission.

CASE 2
An 11 year old girl presented with a two month history of bloody diarrhoea, lower abdominal pain, recurrent mouth ulceration, and a 6 kg weight loss. On examination she was afebrile with a soft but tender abdomen. Investigations showed haemoglobin 85 g/l, WBC 9.0 $\times$ 10$^9$/l, platelets 520 $\times$ 10$^9$/l, ESR 36 mm/h, and C reactive protein 41 mg/l. The presumptive diagnosis was inflammatory bowel disease with active colitis, which was confirmed at colonoscopy. She was transfused and started on intravenous hydrocortisone, antibiotics, and fluids with initial improvement. However, after four days of treatment, she deteriorated with increasing pain, diarrhoea, fever, and abdominal tenderness. An emergency subtotal colectomy was performed with the formation of a mucous fistula and ileostomy. Histology confirmed fulminant ulcerative colitis. Her initial postoperative course was unremarkable, but on the fifth postoperative day she collapsed with dehiscence of the mucous fistula into the peritoneal cavity, resulting in peritonitis. This was corrected surgically but she remained unwell and a pelvic abscess was suspected. Ultrasound showed a fluid collection behind the bladder but could not clearly identify the defunctioned rectum. Computed tomography showed the fluid collection to be the severely inflamed, thick walled rectum distended with fluid. There was an inflammatory mass in the presacral space, with lymphadenopathy along the line of the retained colon (fig 3A and B).

Surgical removal of the rectum was considered but a conservative approach was adopted, with rectally administered steroid enemas and drainage of the rectal fluid. Her condition improved but over the next three weeks she continued to have recurrent fevers and developed deep perianal and sacral ulcers and her inflammatory indices remained persistently high (C reactive protein 110 mg/l). Computed tomography was repeated and showed improvement in the rectal wall with reduced presacral swelling, but there were multiple peri-
anal sinus tracts extending from the presacral space into the gluteal muscles together with dehiscence of the abdominal wound around the fistula. No discrete abscess was identified (fig 4A and B).

With the information provided by the computed tomography, conservative management was pursued with increased nutritional supplementation and a reduction in oral steroids. She was discharged 42 days after surgery on prednisolone enemas. Currently at six months' follow up she is well.

CASE 3
A 3.5 year old girl with Crohn’s colitis diagnosed at age 2 years presented in relapse with severe perianal pain, bloody diarrhoea, and recurrent fever. Investigations showed haemoglobin 82 g/l, WBC 7.8 × 10^9/l, platelets 537 × 10^9/l, ESR 67 mm/h, and C reactive protein 100 mg/l. Examination under anaesthesia showed erythema of the labia and an apparent fistula opening into the left labium. Abdominal x ray showed an abnormal colonic gas pattern consistent with severe colitis. Ultrasound revealed thickened loops of bowel deep in the pelvis but an abscess could not confidently be excluded. The extent of the perianal disease was such that a contrast study or colonoscopy could not be tolerated. Computed tomography was therefore performed to assess her perianal disease further and to exclude a fluid collection. This showed mucosal irregularity of the rectal wall extending into the rectosigmoid, with perirectal sinus tracts and a fistulous connection into the vagina. No abscess was seen.

She was started on enteral nutrition with oral prednisolone and azathioprine, but remained unwell with persistent fevers, abdominal pain, diarrhoea, and raised C reactive protein. Stool was frequently seen at the introitus.

Supine abdominal x ray confirmed mucosal irregularity in the descending colon but there was no evidence of toxic colitis. A limited barium enema under sedation revealed a distended distal colon and rectum, with thickened irregular walls. Ultrasound identified a pelvic mass but it was not possible to determine whether this represented a pelvic collection or distended bowel with thickened wall. Computed tomography was repeated and confirmed markedly thickened sigmoid colon and rectum, with dilatation consistent with colonic Crohn’s disease. In addition, the rectovaginal fistula and a probable rectovesical fistula were demonstrated, neither of which had been visualised by other imaging methods (fig 5A and B).

With this information provided by the computed tomography scan, medical treatment was continued and triple antibiotic therapy...
Table 1  Relative merits of imaging methods in paediatric inflammatory bowel disease

<table>
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<th>Barium studies</th>
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<tr>
<td>Identifying early disease, mucosal oedema and ulceration, enterocutaneous fistulae, and longitudinal extent of disease</td>
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<tr>
<td>Identifying bowel wall changes, mesenteric inflammation and abscess</td>
<td>Identifying bowel wall strictures, mesenteric inflammation, abscesses and fistulae—enterocutaneous, enterovesical, or retroperitoneal</td>
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<td>Demonstration of fistulae (Ultrasound)</td>
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<td>Radiation dose 6.0-9.0 mSV</td>
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(cefotaxime, gentamicin, and metronidazole) added. In view of her severe perianal and rectal disease a defunctioning loop ileostomy was fashioned. Following this, her condition rapidly improved, inflammatory indices improved, and a normal diet was introduced. She remains well at six months' follow up.

Discussion

Barium studies and colonoscopy remain the initial first line diagnostic investigation of a patient with suspected Crohn’s disease. The ability of ultrasound to visualise the bowel wall and its surrounding mesentery is now a valuable diagnostic tool which may have a role in the initial diagnosis of Crohn’s disease and can be used to evaluate the attendant complications. Ultrasound is readily available, flexible, cheap, and has the advantage of not exposing the paediatric patient to ionising radiation. However, in some patients it is difficult to define the full extent of disease with ultrasound, particularly in those patients with abdominal wall tenderness, or in the postoperative period. It can be difficult to exclude confidently the presence of an abscess at ultrasound, and sinus tracts may be difficult to follow. Magnetic resonance imaging (MRI) may specifically look at perianal disease but has limited use in other areas in patients with inflammatory bowel disease. There is limited availability and experience of this technique, and there is no suitable bowel contrast. Many patients find an MRI scan an unpleasant experience.

Computed tomography, especially where spiral scanning is available, provides a rapid, painless, and highly informative investigation which can be accomplished within a few minutes at no discomfort to the patient. With spiral scanning, data can be reconstructed in other planes to provide further useful information.

In the three cases illustrated, computed tomography yielded information regarding the condition of the rectum, identifying intra-abdominal collections, extramural disease, involvement of other organs, and enterovesical and enterocutaneous fistulae. Deep pelvic structures are easily visualised and the ability to obtain good images is not limited by bowel gas or abdominal wall tenderness. This mode of imaging provided information which was not available on ultrasound scans and subsequently determined surgical or medical management.

The use of computed tomography in inflammatory bowel disease is well documented. In one study it led to a change in patient management in 22 of 80 adult patients with Crohn’s disease, and identified unsuspected findings in 13 patients. There are reports of computed tomography in paediatric patients with inflammatory bowel disease, although its use is less well established.

The radiation involved in computed tomography limits its use as a first line investigation. However, the radiation from current scanners from a pelvic examination (8.0 mSV) is the equivalent of a barium follow through or a barium enema and may yield more information in the sick paediatric patient with complicated inflammatory bowel disease (table 1). In our experience computed tomography is of great value if ultrasound fails to resolve the clinical problem.

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