Management of growth failure in Crohn's disease

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Crohn's disease is one of the chronic inflammatory bowel diseases which at present is still of unknown cause. It has been known for some time that this disorder may lead to significant impairment of linear growth and delay in the onset of puberty. Indeed Crohn's disease in childhood may present as short stature or delay in pubertal maturation without any gastrointestinal symptoms. There may be irreversible consequences in terms of reduced potential for height unless this diagnosis is made promptly and treatment instituted to induce remission of disease activity, thus permitting restoration of normal growth. A full blood count, erythrocyte sedimentation rate, and C reactive protein estimation should be standard screening tests for short stature.

In this paper I shall give an account of my own personal practice in the management of children with Crohn's disease and chronic inflammatory bowel disease in the paediatric inflammatory bowel disease clinic which was first established at St Bartholomew's Hospital in 1978 and transferred to the Royal Free Hospital in 1995.

Definitions of growth failure
Growth failure may be defined in various ways: (1) by a static measurement of height below the third centile; (2) by evidence of significant growth faltering as assessed over a period of time by centile chart analysis (that is, a shift from higher to lower centiles of height attained); (3) by showing a reduction in height velocity, which is the most accurate measurement of growth. Measurement of height velocity requires accurate measurement of height over a number of months (a minimum of three months). It may be defined as: (1) height velocity of less than the third centile by centile chart analysis; (2) height velocity < -2.0 SDS (standard deviation score).

For research comparisons, standard deviation scores or Z scores are useful for defining changes in height percentile or in height velocity, but for routine clinical management centile chart analysis is both practical and adequate.

Prevalence of growth failure
A survey of 96 children attending the paediatric inflammatory bowel disease clinic showed that 23% had heights below the third centile but 36% had height velocity below the third centile. The American experience is similar.

This is chiefly a problem of the adolescent and preadolescent years. The height of most younger children with Crohn's disease is normal.

Long term outcome of growth failure
There are three published studies of children with chronic inflammatory bowel disease from prepuberty or early puberty through puberty until cessation of growth. While most patients achieved a final height between the fifth and the 97th centile the distribution was skewed towards the lower centiles, and in some children with Crohn's disease a significant reduction of final height was found. A single observation of 60 adults with so called juvenile onset of chronic inflammatory bowel disease showed that only three had impaired adult growth stature.

The impact of disease on growth would be expected to be less in those patients whose disease preceded their peak growth velocity. As it is known that growth velocity is reduced in children with prepubertal onset of inflammatory bowel disease, it is difficult to compare outcome in such clinical series when populations are not at similar risk.

Clearly advances in modern treatment of chronic inflammatory bowel disease, both medical and surgical, are likely to have a beneficial outcome upon final height.

Pubertal delay and growth failure
Striking pubertal delay may occur in patients in whom a remission has never been successfully induced or in whom relapses have been frequent. In the absence of a remission of disease activity the age of onset of puberty may be delayed indefinitely.

When the age of onset of puberty was taken as breast stage 2 (B2) in girls and testicular volume of 4 ml (TV4) in boys, a significant delay in onset of puberty was found in adolescents with chronic inflammatory bowel disease (fig 1).

Precise pubertal staging (1-5) using the Tanner classification is important, and involves accurate measurement of testicular volume in boys using an orchidometer. Height increase during puberty may be impaired in these children.

There is a paradoxical effect in relation to delayed puberty. It is known that in normal children with similar prepubertal heights late
Pathogenesis of growth failure

The suppression of linear growth that occurs in children with active Crohn’s disease has been attributed chiefly to malnutrition. It has indeed been shown that restoration of nutrition, either by the enteral or the parenteral route, may restore normal growth in these children. However, there is evidence that the inflammatory process per se may itself be a major factor in leading to growth retardation. This evidence includes the observation that products of the activated macrophage such as tumour necrosis factor-α (TNF-α) may be increased in the serum of children with active Crohn’s disease. This increase may synergise with other products of activated macrophages, for example interleukin-1 which may also be raised in Crohn’s disease. Excess production of TNF-α alone, however, caused growth failure and wasting in TNF-α transgenic mice. This could be prevented by an anti-TNF-α monoclonal antibody. Chronic elevation of serum TNF-α or interleukin-2 (IL-2) may have direct inhibitory effects upon new bone formation at the epiphyseal plate.

Monitoring of growth

Accurate measurement of height and plotting on a centile chart, together with calculation of growth velocity, are central to management. A fall in height velocity may be the earliest manifestation of Crohn’s disease. Likewise a subsequent fall of height velocity after a period of successful management may indicate relapse before overt gastrointestinal symptoms occur. The careful documentation of accurate pubertal staging, for example the accurate recording of testicular volumes as discussed above, is also of key importance. Arrest of pubertal development may represent early relapse. It is urgent in these circumstances to induce remission of disease. Radiological determination of bone age may be very helpful in growth monitoring and indicate the prognosis for growth. When there is significant delayed bone age this carries a good prognosis for growth potential.

Close partnership between paediatric gastroenterologist and paediatric endocrinologist is particularly valuable. Joint clinics every four months for children with growth failure and inflammatory bowel have been held in my hospital since 1985. These have been invaluable and educational.

Management of growth failure

In my own practice, enteral nutrition is now the first line treatment for all children with Crohn’s disease, both for small and large intestinal disease, although the evidence for success is better for small intestinal disease. Enteral nutrition may be effective both by restoring nutrition when impaired or inducing a remission of disease activity.

When remission is induced by such a strategy, it carries an excellent prognosis for growth. Some years ago we showed in children that enteral nutrition was as effective as steroids in inducing a remission in Crohn’s disease of the small intestine. When effective, enteral nutrition carries a better outlook than steroids for growth retardation and avoids the risk of growth impairment from steroids per se.

Continued use of nasogastric supplements before completion of puberty has been associated with improved linear growth.

Enteral nutrition may induce a remission by reducing mucosal cytokine production in some at present unknown manner. Both whole protein, that is polymeric, and protein hydrolysate feeds, that is semi elemental, have been shown to be effective in inducing a remission.

Growth hormone deficiency is not a significant factor in most cases. Normal growth hormone secretion has been shown in growth retarded children with Crohn’s disease who were not treated with steroids, although low
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Table 1  Effect of surgery on growth velocity in 40 children grouped according to disease location at time of primary operation

<table>
<thead>
<tr>
<th>Site of disease</th>
<th>No of patients</th>
<th>No &lt; 3rd centile</th>
<th>Mean age at surgery (years)</th>
<th>Growth velocity*</th>
<th>Percentage with increase in growth velocity</th>
<th>Percentage with &gt;10 fold increase in growth velocity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Small bowel</td>
<td>6</td>
<td>2</td>
<td>14.5</td>
<td>2.77</td>
<td>7.28</td>
<td>83</td>
</tr>
<tr>
<td>Ileocecal</td>
<td>20</td>
<td>9</td>
<td>14.3</td>
<td>2.72</td>
<td>6.92</td>
<td>88</td>
</tr>
<tr>
<td>Colonic</td>
<td>14</td>
<td>7</td>
<td>13.4</td>
<td>1.92</td>
<td>7.44</td>
<td>92</td>
</tr>
<tr>
<td>Total</td>
<td>40</td>
<td>18</td>
<td>14.0</td>
<td>2.45</td>
<td>7.18</td>
<td>89</td>
</tr>
</tbody>
</table>

* Growth velocity has been calculated for a minimum of 12 months before and after surgery.

urinary growth hormone secretion was found in patients on steroids. It is well known that steroids may blunt the growth hormone response.

Although thyroid function is typically normal in children with Crohn’s disease, there may be a reduced conversion of T4 to T3 leading to reduced T3 levels. These return promptly to normal on induction of remission.

In boys when there is extreme delay of puberty, that is, testicular volume below 10 ml, monthly intramuscular injections for 3-6 months of testosterone enanthate 125 mg may be helpful, especially when there is clinical remission of disease (Savage MO, personal communication). However, all medical treatment (enteral nutrition and drug therapy) may fail due to either (1) inability to induce or sustain an adequate disease remission, or (2) dependency on a dose of steroids which itself may impair growth, even when steroid sparing treatment with azathioprine has been used.

Surgery may then offer a dramatic opportunity to produce a long standing remission and so restore growth, with the possibility of allowing catch up growth and a normal pubertal growth spurt when correctly timed (table 1; fig 2).

Thus surgery for growth retardation is indicated when there is failure of other treatment to induce an adequate remission. This surgical option is only possible when the disease is not too extensive and it is feasible to resect bowel (table 2).

Right hemicolecotomy, limited small bowel resection, stricturoplasty, and subtotal colectomy with ileostomy are all options, depending on individual circumstances.

When all gross disease can be resected, this has the best prognosis. It is vital to have up to date knowledge of the extent and severity of the disease at the time when surgery is contemplated. This can only be acquired by colonoscopy with ileoscopy and barium follow through studies. Sometimes upper endoscopy, ultrasound, and abdominal computerised axial tomography scans are also required.

Surgical treatment performed up to the early stages of puberty may be dramatically helpful but if delayed until after puberty is well established, the opportunity for growth restoration may be very limited or it may not be possible to achieve it at all (table 2).

Whatever strategy is used to produce a remission, its completeness and duration through puberty is a key factor in influencing growth potential.

Conclusion

Increasing awareness of the importance of slowing of growth and development, both as a mode of presentation of Crohn’s disease in children and as a complication of established disease, should allow greater accuracy in the early diagnosis of Crohn’s disease and in the early recognition of relapse, and enable the most appropriate treatment to be given. When timed appropriately, modern treatments offer the opportunity for children to achieve their full growth potential despite having Crohn’s disease. There is a window of therapeutic opportunity which must be taken advantage of before puberty is complete.

I wish to thank Dr M O Savage who has been my colleague in the joint growth inflammatory bowel disease clinic from 1985 to the present and to the Crohn’s in Childhood Research Association (CICRA) which has supported so much of the clinical research upon which my present clinical practice is based.

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