Jejunal mucosal morphometry in children with and without gut symptoms and in normal adults

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SUMMARY Nineteen diagnostic peroral biopsy specimens from 18 children without diarrhoea, vomiting, or abdominal pain ('control' children) were compared with those taken from 23 children with diarrhoea of varying aetiology to establish the morphometric characteristics of jejunal mucosa in childhood. Comparison was also made with normal jejunal mucosa from adults. Statistical analysis of each characteristic individually showed no significant difference between the 'control' children and those with diarrhoea, but there were significant differences between the mucosae of 'control' children and those of adults: the villi tended to be shorter and the crypts longer in children. Thirty-seven per cent of specimens from the 'control' children showed a partial villous atrophy, that is, they were abnormal by adult criteria. Discriminant analysis of the features measured showed effective separation of the following groups: normal histology from partial villous atrophy in children, healthy adults from 'control' children, and normal histology in adults from normal histology in children.

Morphometric measurements of the jejunal mucosa obtained by peroral biopsy in adults have been exhaustively reported. The techniques employed have varied from simple measurements using linear scales and grids to use of the Weibel template to measure surface-to-volume ratio and television image analysis. There have been fewer reports on jejunal morphometry in children. Jos and other authors used simple micrometer measurements, Risdon and Keeling the surface-to-volume ratio method of Dunnill and Whitehead, and Meinhard et al. computer analysis of projections traced on to computer cards.

The validity of judging children's jejunal mucosae on adult morphological measurements is doubtful. This applies particularly to infants in whom the gut is exposed for the first time not only to postnatal colonisation but also to the influences of food and bacterial antigens which invoke the local immune response. Stereomicroscopic viewing of children's biopsies rarely shows the tall villi seen in normal adult biopsies, and ridges or even convolutions may be seen even when histology does not show gross architectural changes. Minor histological abnormalities are so common that the difference between 'normal' appearances and a 'mild partial villous atrophy' becomes indistinct, cf. We have therefore made simple morphometric measurements on jejunal mucosa to ascertain the following points:

1. Are the measurements in children without gastrointestinal symptoms similar to those of adults?
2. Is the subjective grouping of children's biopsies into normal and partial villous atrophy clinically relevant, or should minor abnormalities be disregarded?
3. Is there a difference between the histological measurements in children without gastrointestinal symptoms and in those with chronic diarrhoea?

Patients and controls

Forty-one patients from whom adequate jejunal biopsies were obtained were referred by paediatricians over a number of years for routine biopsy to exclude small bowel disease when this was diagnostically relevant. This was carried out with the informed consent of the child's parent. The patients were grouped on clinical grounds into those without diarrhoea, vomiting, or abdominal pain at the time of investigation (18 patients) and those with diarrhoea of several weeks' standing (23 patients). The first group of patients (Table 1) is subsequently referred to as
Table 1  'Control' children (without diarrhoea)

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>No. of patients</th>
<th>Range of age at biopsy</th>
<th>No. of biopsies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Failure to thrive</td>
<td>6</td>
<td>10m-7yr 8m</td>
<td>3</td>
</tr>
<tr>
<td>Cow's milk protein intolerance on milkless diet</td>
<td>2</td>
<td>1yr 3m &amp; 2yr 10m</td>
<td>2</td>
</tr>
<tr>
<td>Normal (coeliac disease in family)</td>
<td>3</td>
<td>4yr 4m, 10yr 6m, 12yr 1m</td>
<td>3</td>
</tr>
<tr>
<td>Diarrhoea in infancy (to exclude coeliac disease)</td>
<td>2</td>
<td>3yr 3m &amp; 12yr 6m</td>
<td>1</td>
</tr>
<tr>
<td>Rickets (normal serum Ca++*)</td>
<td>1</td>
<td>2yr 8m</td>
<td>1</td>
</tr>
<tr>
<td>Renal tubular acidosis</td>
<td>1</td>
<td>2yr 4m</td>
<td>1</td>
</tr>
<tr>
<td>Incontinentia pigmenti</td>
<td>1</td>
<td>6yr 7m</td>
<td>1</td>
</tr>
<tr>
<td>Crotul's disease + Still's</td>
<td>1</td>
<td>9yr &amp; 11yr 10m (2 biopsies)</td>
<td>1</td>
</tr>
<tr>
<td>Weight loss? nutritional</td>
<td>1</td>
<td>3yr 6m</td>
<td>0</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>18 + 1 repeat biopsy</strong></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

N = normal.  
PVA = partial villous atrophy.

N = normal. PVA = partial villous atrophy.  
Incontinia pigmenti was subsequently isolated.

'The clinical data of the second group are given in Table 2. Parasites or pathogens were not found in the jejunal juice and/or faeces of the diarrhoea group with the exception of one patient with *Giardia lamblia*. The average age of the 'control' patients was 5 years 5 months whereas that of the children with chronic diarrhoea was 2 years 4 months. The effect of this age difference is discussed later in the statistical analysis but is probably not important. The clinical category as well as the histological appearances of the biopsies had been established by one of us (MS), who was unaware of the morphometric measurements carried out independently by one of the other authors, with checks by two others. All patients with flat biopsies thought to be characteristic of coeliac disease were excluded. One patient in the 'control' group (Table 1) had two jejunal biopsies at an interval of two years and has been included in the statistical analysis as two independent patients. Other patients each had a single biopsy.

The children's biopsies were compared with those taken from seven healthy adult volunteers. In addition, we studied seven biopsies from adult patients which were judged to be normal histologically. The adult groups differed from each other in that the volunteers had deeper crypts, but the values for each group fell within the normal range established in this hospital (68-160 μm). The aggregated adult biopsies are referred to as 'histologically normal'.

Table 2  Children with persistent diarrhoea

<table>
<thead>
<tr>
<th>No. of patients</th>
<th>Range of age at biopsy</th>
<th>No. of biopsies</th>
</tr>
</thead>
<tbody>
<tr>
<td>23*</td>
<td>5m-10yr 6m</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>Mean age 2yr 4m</td>
<td>11</td>
</tr>
</tbody>
</table>

N = normal. PVA = partial villous atrophy.
*Includes one child from whom *Giardia lamblia* was subsequently isolated.

Methods

Only well orientated sections from jejunal biopsies were examined. The morphometric analysis of the sections was performed using a microscope fitted with a Dynascope (model 7A) viewing head. An eyepiece graticule was placed in a ×10 focusing eyepiece, which was fitted with the Dynascope correction collar. The eyepiece graticule scale was calibrated with a stage micrometer. A single division of the graticule scale was found to correspond to 52 μm when used with a ×10 objective and 13 μm when used with a ×40 objective. Subsequently, a 1 cm² eyepiece graticule divided into 100 small squares was fitted into the ×10 focusing eyepiece together with the correction collar. Using a stage micrometer the grid was calibrated and found to have an area of 0·06 mm² when the ×40 objective was selected.

**VILLUS HEIGHT**

Twelve villi were measured in each biopsy from the tip to the villus-crypt junction.

**CRYPT DEPTH**

Twelve crypts were measured in a similar way to the villi.

**ENTEROCYTE HEIGHT**

Ten enterocytes were measured from the middle portion of villi. Only one cell was measured from the side of any villus. Our observations confirm the experience of others that variations in cell height make it difficult to get accurate values with this technique.

**THELIOLYMPHOCYTES (INTRAEPITHELIAL LYMPHOCYTES)**

The number of lymphocytes lying within the epithelium of villi for a total of 500 enterocytes was counted. The lowest quarter of each villus was
excluded from the count. Results were expressed as thelilolymphocytes per 100 enterocytes.

**Inflammatory cells**

These were counted in two areas of the lamina propria: in the villi (villus inflammatory cells) and between the muscularis mucosae and villus-crypt junction (crypt-associated inflammatory cells).

The number of inflammatory cells occurring within 50 small grid squares was counted and expressed as the number of cells per square millimetre of lamina propria.

**Statistical methods**

The significance of differences between the arithmetic means of individual variables (villus height, etc.) was assessed using Student’s t test (two-tailed). Results and discussion using this method are headed ‘Simple analysis’. To determine whether, if all the variables were considered together, the biopsies from different patients or histological categories could be separated into different groups discriminant functions were used.

**Results**

**Simple analysis**

Tables 1 and 2 show the incidence of partial villous atrophy in the control and diarrhoea group of biopsies was 7/19 and 11/23 respectively. This difference is not significant (p = 0.54 exact test).

Table 3 shows the values of the measurements for the seven healthy volunteers, 18 ‘control’ children (one biopsied twice), and 23 children with persistent diarrhoea. There was no significant difference between the mean values of any of these measurements for the ‘control’ and diarrhoea groups of children, but the villus height/crypt depth (VH/CD) ratio and the inflammatory cells (crypt) were near the borderline of significance (p = 0.08 and 0.06 respectively). More detailed analysis (see later) suggested that the significant age difference between the groups was not important. When ‘control’ children were compared with healthy adults significant differences could be shown in villus height, crypt depth, VH/CD, and the number of villus inflammatory cells.

Table 4 compares the measurements for adult and child biopsies graded subjectively as normal (N) or partial villous atrophy (PVA). It is therefore a description in morphometric terms of the histological categories N as applied to children and adults and PVA as applied to children. When the 24 N child biopsies were compared with the 14 adult biopsies the difference in villus height was just statistically significant, but highly significantly different values were obtained for crypt depth and VH/CD. Comparison of the 24 N child biopsies with the 18 PVA child biopsies confirmed the subjective histological grading with significant differences in villus height, enterocyte height, and inflammatory cell count (villi only). The crypt depth measurements showed little difference. A scatter diagram (not presented here) for all the children’s biopsies showed no obvious correlation between villus height and crypt depth, and the highly significant difference in VH/CD must be due to the villus height.

The measurements of these children’s biopsies were also grouped so as to compare the categories N ≥ 0.39

<table>
<thead>
<tr>
<th>Number of biopsies</th>
<th>7</th>
<th>19</th>
<th>23</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age (yr)</td>
<td>46</td>
<td>5yr 5m</td>
<td>0.0042yr 4m</td>
</tr>
<tr>
<td>Villus height (μm)</td>
<td>172</td>
<td>304</td>
<td>0.3</td>
</tr>
<tr>
<td>Crypt depth (μm)</td>
<td>132</td>
<td>157</td>
<td>0.3</td>
</tr>
<tr>
<td>Ratio (VH/CD)</td>
<td>2.83</td>
<td>1.88</td>
<td>0.08</td>
</tr>
<tr>
<td>Enterocyte height</td>
<td>32.3</td>
<td>31.5</td>
<td>0.5</td>
</tr>
<tr>
<td>Theilolymphocytes (villi per mm²)</td>
<td>17.7</td>
<td>19.1</td>
<td>0.2</td>
</tr>
<tr>
<td>Inflammatory cells (villi) per mm²</td>
<td>3100</td>
<td>12700</td>
<td>0.03</td>
</tr>
</tbody>
</table>

**Table 3** Morphometric measurements for patient categories: mean values and p of differences (standard deviations in parentheses)

<table>
<thead>
<tr>
<th>Healthy adult volunteers</th>
<th>‘Control’ children</th>
<th>Diarrhoea children.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of biopsies</td>
<td>7</td>
<td>19</td>
</tr>
<tr>
<td>Mean age (yr)</td>
<td>46 (9yr)</td>
<td>5yr 5m</td>
</tr>
<tr>
<td>Villus height (μm)</td>
<td>172 (48)</td>
<td>304 (52)</td>
</tr>
<tr>
<td>Crypt depth (μm)</td>
<td>132 (13)</td>
<td>157 (38)</td>
</tr>
<tr>
<td>Ratio (VH/CD)</td>
<td>2.83 (0.77</td>
<td>1.88 (0.40)</td>
</tr>
<tr>
<td>Enterocyte height</td>
<td>32.3 (1.6)</td>
<td>31.5 (3.8)</td>
</tr>
<tr>
<td>Theilolymphocytes (villi per mm²)</td>
<td>17.7</td>
<td>19.1</td>
</tr>
<tr>
<td>Inflammatory cells (villi) per mm²</td>
<td>3100</td>
<td>12700</td>
</tr>
</tbody>
</table>

**Table 4** Morphometric measurements for histological categories normal and partial villous atrophy

<table>
<thead>
<tr>
<th>No. of biopsies</th>
<th>14</th>
<th>24</th>
<th>18</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age (yr)</td>
<td>46</td>
<td>16</td>
<td>3</td>
</tr>
<tr>
<td>Villus height (μm)</td>
<td>368</td>
<td>322</td>
<td>0.0001</td>
</tr>
<tr>
<td>Crypt depth (μm)</td>
<td>119</td>
<td>169</td>
<td>0.4</td>
</tr>
<tr>
<td>Ratio (VH/CD)</td>
<td>3.17</td>
<td>2.00</td>
<td>0.0001</td>
</tr>
<tr>
<td>Enterocyte height</td>
<td>33.6</td>
<td>34.1</td>
<td>0.0002</td>
</tr>
<tr>
<td>Theilolymphocytes (villi per mm²)</td>
<td>18.9</td>
<td>19.8</td>
<td>0.03</td>
</tr>
<tr>
<td>Inflammatory cells (villi) per mm²</td>
<td>12000</td>
<td>12700</td>
<td>0.04</td>
</tr>
</tbody>
</table>

VH = villus height; CD = crypt depth.

The p values were obtained using Student’s t test.
and PVA within the 'control' and diarrhoea children, and to compare the 'control' and diarrhoea children within the categories N and PVA. The N category appeared similar in the two patient groups, but the PVA category for the diarrhoea children had shorter villi and more theliolymphocytes than for the control children, though these differences were significant only between 5% and 10%. In the diarrhoea (but not the 'control') group, PVA tended to be found in the younger patients.

DISCRIMINANT ANALYSIS

In order to show whether simultaneous consideration of all measurements gave better discrimination into groups the data were analysed by discriminant analysis. The inflammatory cell counts used in the adult groups were those for the villi only.

This statistical technique has become readily available only with the use of computers and is therefore still not generally familiar. The computer, suitably programmed, constructs mathematical expressions (the canonical variables) of the form $ax_1 + bx_2 + \ldots$, where $x_1$, $x_2$ are the different measurements (in this instance villus height, crypt depth, and so on). The coefficients $a$, $b$, \ldots are calculated so as to weight the contributions of the different measurements, the requirement being that the canonical variable should separate the biopsies as well as possible into groups. We consider this to be an important approach to handling the results of histological measurements. It is only rarely that a single measurement can be used to determine a diagnosis, and the canonical variable provides a way of combining all the measurements together, each being given its appropriate weighting.

In this study we have not applied the canonical variables to fresh data, and it must be emphasised that they are likely to work less well on such data than on the data from which they are derived, particularly as the samples we have analysed are small. A discussion of the use of this method in a medical context is given by Fraser et al.25

Analysis of the 42 children’s biopsies showed that the morphometric studies gave a separation into groups (with some overlap) corresponding to the classification of N and PVA (Fig. 1, horizontal scale). A dividing line at $-2.30$ would have 4 Ns to the right and 2 PVAs to the left of it, but the misclassified ones are all close to the dividing line. This was mainly due to measurement of villus height (mean of 332 $\mu$m in N, 237 $\mu$m in PVA; coefficient $-0.89$). Villus height measurements did not provide a firm classification (ranges: N, 259-411; PVA, 110-328), but a dividing point at 285 $\mu$m would classify only 3 PVAs among the Ns, and 4 Ns among the PVAs, that is, only one more biopsy was misclassified than when discriminant analysis was used. However, only three of the six biopsies misclassified by the discriminant analysis were also misclassified using villus height alone. Since the canonical variable uses all available information, it seems likely that it gives the better result. Crypt depth (coefficient $0.41$) made the next biggest contribution, but neither VH/CD and still less crypt depth alone gave good discrimination between N and PVA.

In summary, the canonical variable gave good, though not complete, separation of N from PVA. Villus height measurements alone did nearly as well, but other measurements, including VH/CD, did not.

The separation into 'with diarrhoea' or 'without diarrhoea' ('control') could not satisfactorily be brought about by discriminant analysis any more than by the simpler analysis discussed earlier. If in Fig. 1 a discriminatory line were drawn at 5.70 on the vertical scale, there would be five 'control' patients above it and six diarrhoea patients below. In contrast to the separation for N and PVA, the misclassified patients were often a long way from their appropriate group. This partial separation was due to age of patient, crypt inflammatory cells, and theliolymphocytes, in that order (coefficients $-0.89$, $+0.61$, and $+0.51$ respectively). The largest correlation with age in the 'within groups' correlation matrix was $-0.35$ with crypt depth. The age contribution to the
separation is therefore probably not very important. Also the significant difference in ages between the 'control' and diarrhoea groups in Table 3 is unlikely to have affected the analysis of the results given in that table. There were only two correlations greater than 0.35, namely, villus inflammatory cells with crypt inflammatory cells (0.44) and villus height with enterocyte height (0.37), both positive.

Two comparisons were made between adults and children. The first was to compare the seven healthy adult volunteers with the 19 'control' children. In this comparison, only moderately good separation was found (Fig. 2). This was due to villus height (coefficient -0.85) and crypt depth (coefficient +0.71) in similar amounts. A VH/CD ratio of, say, 2.40 did nearly as well as the canonical variable in separating the groups. Examination of the within-group correlation matrix showed that although as a group the children had shorter villi and deeper crypts than the adults, within the groups there was a positive (though weak) correlation between villus height and crypt depth of 0.31 (that is, longer villi tended to be associated with deeper crypts). A weak positive correlation of villus height with crypt depth was found within groups in all the canonical analyses performed.

In the second comparison, histologically normal jejunal biopsies from the 'control' group were compared with histologically 'normal' adult tissues. The comparison was made using the biopsies from the seven healthy adult volunteers (Fig. 3) and also all 14 histologically normal adult biopsies (Fig. 4). Using all 14 adults (Fig. 4), a discrimination point at 1 would allot one adult to the child group, and vice versa. The analysis showed that separation was very largely due to crypt depth (coefficient +1.00) with some contribution from villus height (coefficient -0.49). No useful separation could be brought about using single measurements. The highest within-group correlation was -0.41, giving the expected effect that more inflammatory cells were associated with shorter villi.

![Comparison of healthy volunteer adults with 'control' children.](image)

![Comparison of histologically normal volunteer adults with histologically normal 'control' children.](image)

![Comparison of histologically normal adults with histologically normal 'control' children.](image)

**Discussion**

The data presented here are in broad general agreement with previously published results. A study of the extensive literature on adults, shows that there is considerable variation between different studies. This could be due to either variations in the populations studied or variations in technique.

There are few studies on children with which our results can be compared. Jos studied biopsies taken from 14 healthy children aged between 8 months and 16 years. His figures were: villus height 400 μm (range 345-515), enterocyte height 34 μm (range 31-38). Our 'control' children gave a mean of 304 μm for villus height and 32-3 μm for enterocyte height. Lyngkaran and others regarded a VH/CD ratio of 3:1 or more as normal in children, but in none of our children was this value reached: the highest VH/CD ratio was 2:7:1. We know of no previous published data on crypt depth in children. A number of reports on thelolymlphocyte counts were comparable with ours, but a recent study shows that caution is needed in interpreting these as they are affected by the choice of denominator against which they are measured.

The purpose of this study was to examine the concept of 'histological normality' in the jejunal biopsies of children. For ethical reasons we were not able to biopsy entirely healthy children, but assuming that our 'control' group is representative of children without gut involvement, it is clear that there are important differences between biopsies of healthy adults and healthy children. Thus, in the latter, the villi were shorter and the crypts deeper, causing a
highly significant difference in VH/CD ratios. There were also increased inflammatory cell counts in children, significant at the 5% level. In all these measurements there was substantial overlap, but 30 out of the 42 (71%) crypt depths for children lay outside the adult range, as against 19 (45%) of the villus height measurements. Thus a proportion of the biopsies from 'healthy' children were abnormal by adult criteria.

There are two approaches when considering the differences between adults and children. Either the 'normal' small intestine in a child is different from that in an adult simply because of age differences, or a proportion at least of 'clinically normal' children have an abnormal small intestine (due perhaps to an increased incidence of gut infections or transient food intolerances). Thus the situation in the child may be analogous to the minor gut abnormalities seen in healthy adults resident in the tropics.32 33

The morphometric definition of N or PVA could readily be achieved by discriminant analysis. Overlap between patient categories often makes division into groups by any single variable impossible. In this study it was possible to translate histological experience of subjectively classifying children's jejunal biopsies as N or PVA into a single canonical variable, incorporating data from measurement of villus height, crypt depth, enterocyte height, and thiolymphiocyte and inflammatory cell counts. That it was possible to do this shows that the data were not grouped at random. However, the boundaries might not have been correctly drawn. That they were correctly drawn is suggested by a previous study34 in which a highly significant increase (p < 0.001) of IgA plasma cell counts was found in PVA in children. Rather than extending the category of 'normal histological appearance' in children, it is better to say that about 40% of healthy children do not have a histologically 'normal' (N) biopsy.

The subjective grading of children's jejunal biopsies is fairly reliable provided the pathologist has considerable experience with mucosal biopsies35 36 and the orientation of the specimen is correct.37 For those with less experience, morphometric measurements can be useful but the 'norm' must be established separately in each histological department. Morphometry in this study has helped to establish the difference between the 'norm' of adults and that of children and also the fact that a partial villous atrophy in children is found in the absence of gastrointestinal symptoms and, conversely, a normal mucosa in children can be associated with the presence of diarrhoea.

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References
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