Intestinal permeability in kwashiorkor

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Abstract
Intestinal permeability can be assessed non-invasively using the lactulose-rhamnose (L-R) test, which is a reliable measure of small intestinal integrity.

Aims—To determine risk factors for abnormal intestinal permeability in kwashiorkor, and to measure changes in L-R ratios with inpatient rehabilitation.

Design—A case-control study of 149 kwashiorkor cases and 45 hospital controls. The L-R test was adapted to study kwashiorkor in Malawi, with testing at weekly intervals during nutritional rehabilitation. Urine sugars were measured by thin layer chromatography in London.

Results—The initial geometric mean L-R ratios (×100) (with 95% confidence interval) in kwashiorkor were 17.3 (15.0 to 19.8) compared with 7.0 (5.6 to 8.7) for controls. Normal ratios are <5, so the high ratios in controls indicate tropical enteropathy syndrome. Abnormal permeability in kwashiorkor was associated with death, oliguria, sepsis, diarrhoea, wasting and young age. Diarrhoea and death were associated with both decreased l-rhamnose absorption (diminished absorptive surface area) and increased lactulose permeation (impaired barrier function) whereas nutritional wasting affected only l-rhamnose absorption. Despite clinical recovery, mean L-R ratios improved little on treatment, with mean weekly ratios of 16.3 (14.0 to 19.0), 13.3 (11.1 to 15.9) and 14.4 (11.0 to 18.8).

Conclusion—Abnormal intestinal permeability in kwashiorkor correlates with disease severity, and improves only slowly with nutritional rehabilitation.

Keywords: kwashiorkor; intestinal permeability, Africa.

Patients and methods

Patients
The study was carried out at Queen Elizabeth Central Hospital in Blantyre, Malawi between January and September 1995. Children admitted with kwashiorkor (including marasmic kwashiorkor) during the study period were eligible for entry into the study. Kwashiorkor was diagnosed clinically on the basis of nutritional oedema, with other causes of oedema such as nephrosis, nephritis, and severe malarial anaemia excluded. The experimental group of 149 children for permeability studies was selected from a total of 606 kwashiorkor admissions during the study period (24.6%) of whom 553 remained in hospital for at least five days. In
selecting patients for a permeability study, there was an explicit bias to boys (from whom it is easier to collect urine), to those admitted just before the two study nights per week, and to those likely to survive for repeat testing. These selection criteria were applied consistently throughout the study period by the same research assistant. Children with diarrhea were rehydrated before permeability testing. Controls were selected from hospitalised children without diarrhea or wasting, who were recovering from pneumonia, soft tissue infections, or chronic diseases such as tuberculosis or osteomyelitis. Children with other diseases or treatments which are known to affect permeability were excluded from the study (for example severe malaria, iron deficiency anaemia, non-steroidal anti-inflammatory drugs).

A parent or guardian of children in the study was administered a field tested questionnaire about household water supply, sanitation, hygiene practices, health, and socioeconomic status. This was developed into a score which increased proportionately with higher socioeconomic status. Weights and recumbent lengths of subjects were measured by a single trained research assistant using standard techniques on a Salter hanging scale and locally made stadiometer. Although measurements were recorded to the nearest 0.1 kg and 0.1 cm, we found from repeat measurements that there was digit preference and a tolerance level closer to 0.5 cm for height. Since all children had recumbent length measured, because it is more reproducible in sick children, a correction factor of 1.5 cm was subtracted from the length measurement of children >24 months to approximate height for the National Center for Health Statistics (NCHS) standard.14

After standard treatment, all kwashiorkor cases were given routine antibiotics (cotrimoxazole) for at least five days, and supplements of minerals, micronutrients, and vitamins. Out of the 149 children with kwashiorkor in whom permeability testing was done, 76 were treated with a milk based diet, 64 with an exclusively maize based (milk free) diet, and nine with both diets over a changeover period. Details of the diets are given in the companion paper.15 Clinical sepsis was defined as fever, respiratory distress, a change in mental status, shock or any abrupt deterioration in condition. Consent for enzyme linked immunosorbent assay (ELISA) testing for HIV infection after precounselling was often refused, making it impossible to test all malnourished children. Although there were only 23 positive tests out of 50, our estimated prevalence of HIV infection in study patients from a clinical protocol was 43 children (29%). HIV infected patients were not excluded from the study because, like tuberculosis and chronic diarrhoea, it is part of the clinical spectrum of kwashiorkor in Africa.

Table 1 Baseline comparison of malnutrition cases and hospital controls

<table>
<thead>
<tr>
<th>Mean (95% CI) or No (%)</th>
<th>Kwashiorkor cases (n=149)</th>
<th>Controls (n=45)</th>
<th>ANOVA p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (months)*</td>
<td>28.3 (26.7 to 29.9)</td>
<td>28.6 (22.5 to 36.3)</td>
<td>0.88</td>
</tr>
<tr>
<td>Male sex</td>
<td>116 (78)</td>
<td>23 (51)</td>
<td>0.001</td>
</tr>
<tr>
<td>Distance (min) to hospital</td>
<td>60.3 (54.1 to 66.8)</td>
<td>59.8 (42.5 to 84.3)</td>
<td>0.98</td>
</tr>
<tr>
<td>Family size</td>
<td>4.7 (4.4 to 5.0)</td>
<td>5.2 (4.7 to 5.9)</td>
<td>0.07</td>
</tr>
<tr>
<td>Still breast fed</td>
<td>8 (5.4)</td>
<td>16 (40)</td>
<td>0.000001</td>
</tr>
<tr>
<td>Socioeconomic status score†</td>
<td>14.8 (14.1 to 15.5)</td>
<td>17.5 (15.8 to 19.1)</td>
<td>0.001</td>
</tr>
<tr>
<td>Mothers’ schooling (years)</td>
<td>3.3 (2.9 to 3.8)</td>
<td>4.0 (2.6 to 6.4)</td>
<td>0.21</td>
</tr>
<tr>
<td>Fathers’ schooling (years)</td>
<td>6.2 (5.6 to 6.9)</td>
<td>7.6 (6.6 to 8.9)</td>
<td>0.06</td>
</tr>
<tr>
<td>Wasting (WHZ)</td>
<td>−1.98 (−1.80 to −2.15)</td>
<td>−0.18 (0.20 to −0.57)</td>
<td>0.00000001</td>
</tr>
<tr>
<td>Stunting (HAZ)</td>
<td>−3.52 (−2.94 to −3.73)</td>
<td>−2.38 (−1.98 to −2.78)</td>
<td>0.001</td>
</tr>
<tr>
<td>Urine volume for L-R test (ml)*</td>
<td>65.3 (58.1 to 73.5)</td>
<td>90.8 (81.8 to 114.6)</td>
<td>0.001</td>
</tr>
<tr>
<td>Mean L-rhamnose recovery (%)</td>
<td>1.04 (0.85 to 1.28)</td>
<td>2.54 (1.85 to 3.49)</td>
<td>0.0002</td>
</tr>
<tr>
<td>Mean lactulose recovery (%)</td>
<td>0.164 (0.145 to 0.187)</td>
<td>0.162 (0.114 to 0.231)</td>
<td>0.95</td>
</tr>
<tr>
<td>L-R ratio*</td>
<td>17.3 (15.0 to 19.8)</td>
<td>7.0 (5.6 to 8.7)</td>
<td>0.000001</td>
</tr>
</tbody>
</table>

* Geometric means.
† Higher score for higher socioeconomic status.
ANOVA = analysis of variance; HAZ = height/age z score; WHZ = height/weight z score.
## Results

We carried out 347 L-R permeability tests in 194 hospitalised children, including 149 with kwashiorkor and 45 controls. Repeat testing was done a week later in 109 kwashiorkor cases (73.2%) and another 44 tests on the third or fourth week. Table 1 compares kwashiorkor cases to controls on admission, showing expected differences in sex, breast feeding, socioeconomic status, and nutritional status. Although there was a male predominance (2.5:1) of kwashiorkor cases studied, mean L-R ratios in boys were 15.8 (14.1 to 17.9) compared with girls 15.1 (12.2 to 18.7). L-R ratios were paradoxically higher for children from families in the top 20% of socioeconomic status (23.9, 18.5 to 31.0) and with over 8 years of maternal schooling (22.4, 14.8 to 36.9), but these variables were not significant on multiple regression of L-R ratios.

Mean L-R ratios (95% CI) for all 149 kwashiorkor cases on admission was 17.3 (15.0 to 19.8) compared with 7.0 (5.6 to 8.7) for controls (p<0.000001). Although 32 kwashiorkor cases (21.5%) had ‘normal’ permeability ratios for this population (5.6 to 8.7) on admission, the repeat ratio after one week of treatment deteriorated in half of them to a mean of 17.1 (14.6 to 20.1). A history of persistent diarrhoea before admission was elicited in 33.8% of kwashiorkor cases, but did not affect mean L-R ratios on admission compared with those without such a history (17.1 vs 17.7). In contrast, an increased duration or severity of diarrhoea in hospital was associated with worse permeability ratios (fig 1). Mean L-R ratios for kwashiorkor complicated by clinical sepsis were higher (18.2 v 13.0, p=0.009), and remained significant when controlled for diarrhoea (F=7.1, p<0.005). A multiple regression model for intestinal permeability (F=24.5, p<0.005), which explained 33.6% of the variance, had the following significant independent variables (partial F tests) for higher L-R ratios: oliguria (31.5), severe diarrhoea (17.4), death (16.4), treatment with a maize diet (11.0), young age (10.1), and wasting (4.1).

Differences in mean L-R ratios between kwashiorkor and controls were due to higher L-rhamnose absorption in controls. But in kwashiorkor, the combination of increased lactulose permeation and decreased L-rhamnose absorption was associated with a higher risk of

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### Table 2 Logistic regression for death and diarrhoea

<table>
<thead>
<tr>
<th>Dependent variable</th>
<th>Independent variable</th>
<th>Odds ratio</th>
<th>95% CI</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Death</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Rhamnose absorption*</td>
<td>0.04</td>
<td>0.01 to 0.15</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td></td>
<td>Lactulose permeation*</td>
<td>17.4</td>
<td>5.4 to 56.1</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td></td>
<td>Maize diet (v milk†)</td>
<td>3.7</td>
<td>1.6 to 8.3</td>
<td>0.002</td>
</tr>
<tr>
<td></td>
<td>Younger age*</td>
<td>16.1</td>
<td>2.2 to 118.4</td>
<td>0.006</td>
</tr>
<tr>
<td></td>
<td>Diarrhoeal burden*</td>
<td>2.8</td>
<td>1.2 to 6.9</td>
<td>0.021</td>
</tr>
<tr>
<td></td>
<td>Rash severity</td>
<td>1.6</td>
<td>1.0 to 2.4</td>
<td>0.031</td>
</tr>
<tr>
<td></td>
<td>Wasting severity (WHZ)</td>
<td>1.5</td>
<td>1.0 to 2.2</td>
<td>0.045</td>
</tr>
<tr>
<td>2. Diarrhoea</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Rhamnose absorption*</td>
<td>0.27</td>
<td>0.2 to 0.9</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td></td>
<td>Lactulose permeation*</td>
<td>5.1</td>
<td>2.4 to 10.7</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td></td>
<td>Milk diet (v maize†)</td>
<td>4.8</td>
<td>2.7 to 8.6</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td></td>
<td>Oedema severity</td>
<td>1.6</td>
<td>1.3 to 2.0</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>

* Log values.
† See companion paper for explanation of diets.
Diarrhoeal burden = proportion of hospital days with diarrhoea.
WHZ = weight/height z scores.
Intestinal permeability in kwashiorkor

Figure 2  The higher probability of death with decreasing L-rhamnose absorption at different lactulose permeation rates. Note the logarithmic scale for lactulose and L-rhamnose. This illustrates the inverse ratio of L-rhamnose and lactulose permeation in kwashiorkor, which is consistent with villous atrophy.

Discussion
PERMEABILITY IN KWASHIORKOR

In this first study to focus on intestinal permeability in kwashiorkor, we found permeability ratios of >10 on admission in 110 kwashiorkor cases (73.2%), indicative of mucosal damage. Kwashiorkor is known from biopsy studies to be associated with villous atrophy, decreased villous-crypt ratio and increased cellularity of the lamina propria. Although severe mucosal injury occurs in only a proportion of malnourished children, it is more common in kwashiorkor, possibly due to the effect of protein depletion on mucosal recovery. An Ethiopian study of 17 children with kwashiorkor documented abnormal D-xylose tolerance and carbohydrate malabsorption in a high proportion of cases. The mean L-R ratios in our kwashiorkor subjects were still lower than those reported for acute gastroenteritis (43, CI 26 to 60) but higher than for persistent diarrhoea in the UK (12, 8 to 16) by the same technique. The explanation for the rise in L-R ratio in those with low ratios on admission is unclear, but it corresponded to nosocomial infection and clinical deterioration.

Malawian control children also had abnormal mean L-R ratios (7.0, 5.6 to 8.7) compared with UK control children (2.7, 0.8 to 5.2) using the same test solution and laboratory. Only eight cases of kwashiorkor (5.4%), and nine controls (20%) in this study had L-R ratios in the normal range for UK children. This is a feature of the tropical enteropathy syndrome, with small intestinal mucosal damage from overexposure to enteric pathogens in a contaminated environment, often with small bowel bacterial overgrowth. A study in beagles found small intestinal bacterial overgrowth was associated with abnormal permeability in spite of normal gut histology. Permeability testing is obviously a more sensitive indicator of subtle or patchy mucosal abnormalities than morphometry. The persisting abnormal permeability after clinical recovery in our kwashiorkor cases cannot be explained as tropical enteropathy syndrome, since mean ratios were much higher (14.4 vs 7.0).

A notable feature of intestinal permeability in the present study was its prognostic value in kwashiorkor. Both decreased L-rhamnose recovery (malabsorption) and increased lactulose recovery (impaired barrier function) were independent predictors of mortality on logistic regression (table 2). Malabsorption of L-rhamnose with increased lactulose permeation is typical of the permeability changes with villous atrophy. The main permeability difference between kwashiorkor cases and controls in this study was decreased mean L-rhamnose absorption (table 1), which is thought to reflect villous shortening with loss of absorptive surface due to malnutrition. This is confirmed by the strong correlation (p=0.005) between the severity of L-rhamnose malabsorption and wasting (weight/height z scores).

Abnormal permeability was also associated with three severity of illness factors: oliguria, sepsis, and diarrhoeal severity. Oliguria was measured as urine output (in ml/kg/hour) during overnight testing. Figure 3 shows the rise in L-R ratios at low urine outputs. This was not just a technical problem with the test, but reflected the degree of anorexia due to severe illness. Children with a low urine output during testing (oliguria) had a significantly higher mortality, longer hospital stay and more severe rash. Oliguria was not from dehydration, and was not associated with diarrhoea on the day of testing (p=0.7). Nor was it due to incomplete urine collection because we developed a urine collection protocol (after 12 months of failure) which made incomplete collection unlikely. Moreover, oliguria was associated with a higher urine urea concentration (41.9 vs 26.4 mmol/l,
PERMEABILITY TESTING

The dual sugar test is ideally suited for assessing the small intestine in children with severe malnutrition and diarrhoea, and has important advantages of non-invasiveness, repeatability, and reliability. Malnutrition may affect intestinal function by a decrease in absorptive surface area without decreasing villous height or causing other structural changes.38

The main problem with permeability testing on urine in sick young children, which is not overcome by using two probes, is that they cannot be relied upon to void at the end of a five hour urine collection, so the timed collection would in fact be shorter than the prescribed five hours. Permeation rates of test sugars differ between jejunum and ileum, so urine collections shorter than five hours would give falsely low ratios.39 Although urine collection difficulties in children are well known, this issue has been largely ignored in paediatric permeability studies, possibly because most subjects in developed countries have been older children. We overcame the problems of prolonged urine collection by using overnight collection with urine bags and a special skin adhesive technique. It is unfortunate that the many different methods of dual sugar testing make it difficult to compare results with test sugars of different composition, proportions, or osmolality.40

Finally, in spite of clinical recovery, mean L-R ratios did not return to control values in this study. Only 14 of 44 L-R tests (31.8%) had fallen to a ratio of <10 after 3–4 weeks of nutritional rehabilitation (fig 4). A slow response of the small intestine morphology to nutritional rehabilitation has been noted previously.21–23 41

It has been attributed to persisting exposure to intestinal pathogens due to poor hygienic conditions, but our permeability ratios after clinical recovery are much higher than in tropic enteropathy syndrome as seen in controls.38–41 The high rates of nosocomial diarrhoea, including its presence on the day of testing, its severity (frequency/day) and its duration in hospital, and associated both lactulose and rhamnose permeation. This is consistent with the view that nosocomial diarrhoea in malnutrition affects both mucosal barrier and absorptive functions, whereas malnutrition itself (wasting) affects mainly absorption.

The abnormal permeability associated with clinical sepsis in this study is notable because of studies on bacterial translocation. Abnormal intestinal permeability is a feature of bacterial translocation in malnourished adult patients with multiorgan failure secondary to trauma, sepsis or burns.20–33 The high rates of nosocomial sepsis in kwashiorkor36–38 are consistent with evidence that protein malnutrition predisposes to gut origin sepsis.39 This offers the hope that dietary or other treatment measures which improve gut permeability in kwashiorkor may lower sepsis and case fatality rates.

The relationships between intestinal permeability, AIDS and socioeconomic status in our kwashiorkor cases were puzzling. Mean L-R ratios for the 50 ELISA tested cases were 18.2 (13.3 to 24.9) for positives compared with 24.0 (18.0 to 32.1) for negatives. The remaining 99 untested cases had L-R ratios of 18.2 (9.9 to 33.5) for those with probable AIDS by the clinical protocol compared with 15.4 (12.8 to 18.4) for those not suspected of AIDS. We interpret this to mean that clinical status or severity of disease predicted abnormal permeability better than HIV status, since the difference between tested negative and untested unsuspected cases reflected the degree of sickness and response to nutritional rehabilitation.

Mean socioeconomic scores for AIDS cases with kwashiorkor was 16.0 (4.8 to 17.3) compared with 14.4 (3.5 to 15.2) for negative cases, with no difference whether diagnosed by ELISA test or clinical protocol. The paradox of higher mean L-R ratios in kwashiorkor with parents of higher socioeconomic and educational levels (p<0.005) is explained by AIDS, since it was clinically obvious that AIDS was a more likely cause of malnutrition in these families than in those living in dire poverty.
Key messages

- Kwashiorkor still has a 30% case fatality rate at urban hospitals in Malawi
- Intestinal permeability can be measured non-invasively by the lactulose-rhamnose test as an indicator of gut damage
- The degree of abnormal permeability in kwashiorkor reflects the severity of illness and improves only slowly with nutritional rehabilitation
- Changes in intestinal permeability can be used to evaluate treatment interventions

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