Impact of intestinal permeability, inflammation status and parasitic infections on infant growth faltering in rural Bangladesh

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A longitudinal study of 298 rural Bangladeshi infants found evidence of growth faltering starting at 3 months of age. Anthropometric status declined substantially in the first 2 years of life, with weight-for-height (WHZ) falling from −0·49 to −1·75, weight-for-age (WAZ) from −1·18 to −2·87 and height-for-age (HAZ) from −1·00 to −1·88. Higher concentrations of the acute-phase protein α-1-acid glycoprotein (AGP) and higher gut mucosal damage (as signified by raised lactulose:mannitol (L:M) ratios) were both associated with chronic malnutrition as indicated by poorer HAZ and WAZ scores (P=0·011 and 0·005 for AGP and 0·039 and 0·019 for L:M ratio, respectively). Higher Hb levels were related to improved z-scores, while elevation of Giardia-specific IgM titre (GSIgM) was associated with poor WAZ and WHZ (P=0·015 and 0·039, respectively). IgG did not show any significant association with z-scores and the L:M ratio did not correlate with any of the inflammation markers or Giardia infection. The prevalence of geohelminth infections was low (only 4% in the total study period). However, the level of GSIgM indicated high endemicity of Giardia infection from early in life, although very few cysts were detected from stool samples. These findings suggest that rural Bangladeshi infants are being exposed to high levels of infection with comcomitant gut damage and growth faltering.

Infant growth: Intestinal permeability; Giardia intestinalis; Giardia-specific IgM antibody: α-1-Acid glycoprotein

The burden of undernutrition among most young Bangladeshi children remains very high and a recent nationwide study(3) found that 40·8% of children under 5 years old were stunted, 47·1% were underweight and 11·9% were wasted. These high burdens reflect both shorter- and longer-term insults. Growth faltering which is common in many developing countries starts at an early age with weight-for-age z-scores (WAZ) and weight-for-height z-scores (WHZ) falling from about 3 months of age until about 12 months of age, whereas the decline in height-for-age z-score (HAZ) starts immediately and continues until about 24 months of age(2). The time period between 3 and 24 months of age of a child has been described as ‘the window of opportunity for addressing undernutrition(12,3) and so interventions to improve nutritional status throughout the growth period must be accomplished within this rather short period of childhood.

A hypothesis of a cause of infant growth faltering in developing countries was suggested from work in The Gambia. Lunn(4) and Campbell et al.(5) found that impaired small-intestinal mucosal function explained up to 64% of the observed height and weight faltering of Gambian infants aged between 2 and 15 months of age. They proposed that frequent gastrointestinal infections, arising from unhygienic food preparation or storage, damaged the small-intestinal mucosal function in two ways. First, by the loss of vulnerable brush-border enzymes including lactase, which impaired digestion and absorption, and second by disruption of the barrier function, which allowed translocation of macromolecules across the mucosa and into the body. Although the exact causes of the enteropathy remain unknown, the onset of mucosal damage in these Gambian infants started at about 6 months of age, which coincided with the time of introduction of home-prepared weaning foods as well as the onset of growth faltering.

Parasitic infections due to Giardia intestinalis and geohelminths are very common in many tropical environments, and these infections have been proposed as one major cause of intestinal mucosal damage(6–9). Giardia infection is associated with poor nutritional status of young children in developing countries(6,10) and geohelminths have also been implicated in restricting child growth(11) and are known to cause maladaptation and malassorption(12). It is possible that even low intensities of infections with these parasites have marked effects on the fast-growing infant, and Trichuris trichiura has been shown to be associated with poor growth in both weight and height in Nicaraguan children less than 24 months of age(13).

In Bangladesh, although a number of studies have examined the effect of gastrointestinal parasite infection on the growth of infants, there are little available data on infants and young children. It was surprising that so little attention had been paid by researchers to the role of Giardia infection in growth faltering. It is possible that parasites may have been underestimated as a cause of growth faltering. The data from the current study strongly suggest that Giardia infection is an important cause of growth faltering in rural Bangladesh.

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Abbreviations: AGP, α-1-acid glycoprotein; GSIgM, Giardia-specific IgM titre; HAZ, height-for-age z-score; L:M, lactulose:mannitol ratio; WAZ, weight-for-age z-score; WHZ, weight-for-height z-score.

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children below 24 months of age, although a WHO study on rural Bangladeshi infants showed that 50 % of 2-year-old children had *Ascaris* infection, 25 % *Trichuris* and 21 % had hookworms (CGN Mascie-Taylor, personal communication). Additionally, a cross-sectional pilot survey conducted by the authors in rural Bangladesh found that 87 % of 2- to 18-month-old infants had elevated *Giardia* antibody titres.

The present study describes the changes in anthropometric and biochemical variables over the first 2 years of life of Bangladeshi infants and examines the inter-relationships of infant growth, *Giardia* infection, immune response and intestinal mucosal function.

**Subjects and methods**

The data came from a longitudinal randomised controlled study which examined the impact of regular anti-*Giardia* and anthelmintic treatments on growth, intestinal mucosal function and immune status of infants in a rural area of Bangladesh\(^{16}\). The study, however, generally failed to show any impacts of the treatments, due to the unexpectedly high rate of *Giardia* re-infection. The 36-week trial (June 2003 to April 2004) was carried out in Dhamrai Upazila, a typical rural area located 40 km north-west of Dhaka, the capital of Bangladesh. All infants under 11 months of age in the area were recruited (their age ranged from 3 to 15 months at the beginning of the study) using a list of the national immunisation programme as well as local community knowledge.

After the medical doctor and the trained female local assistants explained the study purposes and the process, parental consent was obtained (a consent form was prepared in Bengali) and families were informed that they could withdraw from the study at any time.

The study was based in four contiguous administrative areas (wards) and at the beginning of the study, socio-demographic and economic information was collected from families including family size, number of siblings, religion, number of possessions (cattle, goat, television, bicycle, motorbike and mobile phone), parental background (age, occupation and education) by trained local female assistants. The Bengali questionnaire was pre-tested in the pilot study and checked for accuracy using an independent translator.

In the longitudinal study, infants had their weight and supine length measured every 4 weeks. Every 12 weeks a lactulose:mannitol (L:M) ratio intestinal permeability test was performed, a capillary blood sample was taken to measure Hb, albumin, α-1-acid glycoprotein (AGP), IgG and *Giardia*-specific IgM titre (GS IgM) and a stool sample was collected and examined for the presence of eggs of the three common geohelminths and *Giardia* cysts.

**Anthropometry**

The infant’s weight and supine length were measured using standard techniques\(^{15}\); length was measured to the nearest 0.5 cm using a portable anthropometric board (TALC, St Albans, Herts, UK) and weight was measured with 0.05 kg precision using a digital electric scale (UC-321; A & D Co., Tokyo, Japan). Inter-observer technical error of measurement and the coefficient of reliability (R) were calculated\(^{16}\) at three different times during the study period.

All values of R were more than 0.95, so the quality of measurement was deemed acceptable throughout the study period. Most mothers remembered their infant’s date of birth, as there was less than a year between the birth and their infant’s recruitment into the study. Birth details were validated whenever possible using the infant’s vaccination card.

**Urine collection for testing intestinal permeability**

The solution for testing intestinal permeability contained 200 mg lactulose (Regulose\textsuperscript{®} syrup, lactulose 3.35 g per 5 ml; donated by General Pharmaceuticals Limited, Gazipur, Bangladesh) and 50 mg mannitol (Sigma-Aldrich, Poole, Dorset, UK) per ml water. Infants received 2 ml of the solution per kg body weight using a disposable dropper with a scale so that any solution lost could be estimated and made up to ensure the infant received the correct volume. The solution is quite sweet and pleasant tasting and infants usually swallowed it readily. Mothers were requested not to feed their infants for at least 1 h before dosing. Infants were encouraged to drink water 30 min after taking the sugar solution in order to help urination. A urine specimen collection bag (Paediatric/Non-sterile U-Bag; Hollister Limited, Aurora, ON, Canada) was attached to the infant immediately after dosing. These urine bags are equipped with a drainage tube so that urine can be removed from the bag soon after each urination, thereby keeping the infant as dry as possible and reducing loss. The urine was collected in a container marked with the infant’s identification number and a few drops (about 50–100 μl) of 0.2 % (w/v) chlorhexidine digluconate solution (Sigma-Aldrich) were added as a bacteriostat.

Each collection time was of 3 h duration, except for the first examination when urine was collected for 5 h. If, however, the infant did not urinate within 90 min of taking the sugar solution, then urine collection was continued for 5 h. The collection of urine for 5 h often proved to be a problem for these rural Bangladeshi mothers as they found it difficult to be absent from their house for such a long time, and 3 h was generally about the maximum time they could spare. Akram et al. have shown good agreement between L:M ratios collected over 2 and 6 h\(^{17}\). On completion of the urine collection and after checking for faecal contamination, total volume was measured and a 2 ml well-mixed sample was taken and kept at –20°C before shipment to the UK for analysis. If an infant had diarrhoea or skin infection of the perineal area, the test was postponed, and repeated as soon as the infant had recovered.

**Blood collection and Hb measurement**

A finger-prick technique was used to collect blood samples from the infants. This method gives minimal pain, is less invasive, causes less distress to infants, and provides sufficient blood for analysis (about 100 μl). One finger was cleaned using disinfectant liquid and, after drying completely, the finger was pricked to a depth of 2.25 mm using a spring-loaded disposable lancet (HemoCue Safety Lancet; HemoCue Ltd, Angelholm, Sweden). A maximum of five drops of blood were collected on a Guthrie card (903 Specimen Collection Paper; Schleicher & Schuell Bioscience GmbH, Dassel, Germany). An additional drop was placed in a micro-cuvette.
(HemoCue Blood Hemoglobin Cuvette; HemoCue Ltd) and the Hb concentration was measured immediately using a portable haemoglobinometer (HemoCue; HemoCue Ltd). Blood spots were dried overnight at room temperature, avoiding any insect contact, and then packed individually in a sealed plastic bag and stored at −20°C with desiccant before analysis.

Stool sample collection and analysis

A day before the stool sample collection, a container was distributed to the mothers, and they were instructed on how to collect a stool sample and to make sure the specimen was not contaminated with soil or dirt. Diarrhoeal stool samples were not examined as they are too diluted and another sample was taken after the infant recovered, usually within 1 to 2 d. The collected specimens were taken as soon as possible to the laboratory in the research station where a 10–15 g sample was preserved in 9 ml of 10% formalin and kept at room temperature. Stool samples were examined at the Parasitology Laboratory at the Laboratory Sciences Division of the International Centre for Diarrhoeal Disease Research, Bangladesh (Dhaka, Bangladesh) where the formalin–ether faecal concentration technique (18) was used to measure the prevalence and intensity (eggs per g faeces) of Ascaris lumbricoides, T. trichiura and hookworm (either Ankylostoma duodenalis or Necator americanus) and cysts of G. intestinalis.

Laboratory assays of urine and blood samples

All urine and blood assays were conducted at the Department of Biological Anthropology, University of Cambridge (Cambridge, Cambs, UK). Enzymic assays were used to measure the concentration of mannitol, lactulose and lactose in urine (19,20).

Plasma constituents were eluted from the Guthrie card blood-spots by 24 h incubation in 1·25 ml of a phosphosaline buffer (0·01 M-sodium phosphate, 0·5 M-sodium chloride, pH 7·2) containing 1% Tween 20. The resulting solution, a 20-to 50-fold dilution of the blood, was used directly for the analyses. An immunoturbidimetric technique was used for the albumin assay (antibody and standards supplied by DakoCytomation, Ely, Cambs, UK). Blood concentrations of IgG and AGP were determined using a standard double-sandwich ELISA (IgG antibodies from DakoCytomation and AGP antibodies from Acris Antibodies GmbH, Hidenhausen, Germany; standards were supplied by DakoCytomation). The Giardia-specific IgM ELISA assay used Giardia trophozoite antigen following the method of Goka et al. (21). The Hb concentration in the eluted sample was analysed using the cyanmethaemoglobin method (Randox Haemoglobin Assay Kit, Randox Laboratories Limited, Crumlin, Co. Antrim, UK) and, by comparison with the Hb reading taken at blood collection, was used to estimate the blood volume of the eluted spot. In order to quote plasma concentrations from the blood spot measurements, a conversion factor was calculated from data in which albumin and IgG were assayed in both blood and plasma of eighty-six urban poor Bangladeshi infants. Thus, plasma concentration = blood concentration × 3·14. The CV of all the assays were below 5%. Precision of measurement was one decimal place for Hb and albumin and two decimal places for other biochemical variables.

Statistical methods

Data were analysed using the SPSS for Windows statistical package (versions 10 to 13; SPSS Inc., Chicago, IL, USA). Sequential multiple regression analyses were mainly used. The L:M ratio distribution was skewed and was normalised using a log10 transformation. Curve estimation was used to check for the linear, quadratic and cubic relationships between two continuous variables. Statistical significance was accepted at the 5% level.

HAZ, WAZ and WHZ of the infants were determined using Epi Info 7 (version 3.3.2) based on National Center of Health Statistics year 2000 references, Centers for Disease Control and Prevention (22). Because the anthropometric measurements were not taken at exactly 4-weekly intervals, individual curve estimates (up to fourth-order polynomial) were applied and from the polynomial regressions the weights and supine lengths at 28 d intervals were calculated using SAS (version 8.2; SAS Institute Inc., Cary, NC, USA). The estimated weight and supine length were used to calculate the HAZ, WAZ and WHZ scores at exactly 4-weekly intervals.

GSiGM titres of infants in the present study were compared with values obtained from sixty-seven UK infants (23) (aged 10–20 months) who were assumed not to have been exposed to Giardia. The upper 95% confidence value (mean 0·154 + 2 × 0·086 = 0·326) obtained for the UK children was used as the cut-off point to delineate Giardia-positive from -negative infections in the Bangladesh study.

Ethical approval was obtained from The Bangladesh Medical Research Council.

Results

Out of the initial sample of 410 infants, 298 infants (143 males and 155 females) provided full information comprising anthropometric measurement at 4-weekly intervals (ten measurements) and blood spot, urine and stool examinations (four examinations) at 12-weekly intervals. There were no significant differences in infants’ age, sex, anthropometry and socio-demographic and economic status between the sample (n 298) and those lost to attrition (n 112).

The infants’ families were generally poor; most lived in houses made of mud with a tin roof (93%), a quarter of families did not own any cattle or goats (39%), nearly half of the fathers (46%) worked as daily labourers and only 7% of mothers were in paid work. About half of the fathers and mothers (45 and 42%, respectively) had received no formal education, and one-third of the mothers (29%) were teenagers.

Stool examination showed that the prevalences of geohelminth infections in the infants were low (4% in the total study period) and very few (<0·9%) of Giardia cysts were detected. However, prevalence using an elevation of GSiGM titres showed that 98% of the infants were infected with Giardia over the study period.

Table 1 presents the overall mean and standard deviation of the three z-scores and the six biochemical variables. The anthropometric variables provide evidence of both acute
wasting, low WHZ) and chronic malnutrition (underweight, WAZ and stunting, HAZ) and most of the children were anaemic. IgG, which has a long half-life (23 d; Lydyard et al. (24)), reflects chronic immune stimulation. Plasma concentrations of the acute-phase protein AGP increase 3- or 4-fold within 3 d of infection and in the absence of further infection the half-life is about 5.5 d (25), so the raised AGP mean value indicates regular re-infection as do the elevated *Giardia* titres; high L:M ratios imply damage to intestinal mucosal function.

All *z*-scores showed a significant downward relationship with age (all \( P<0.001 \); Fig. 1), with HAZ falling nearly 1 SD from \(-1.00\) to \(-1.88\), WAZ and WHZ falling more than 1.5 SD; from \(-1.18\) to \(-2.87\) for WAZ and from \(-0.49\) to \(-1.75\) for WHZ. On average, female HAZ values were consistently higher by about one-third of an SD than males, while WHZ was higher in males (0.22) on average, but the difference became smaller as the infants aged. No sex difference in WAZ was found.

Hb showed a slight upward relationship with age (\( y = 98.393 + (-0.862) \times \text{age} + 0.040 \times \text{age}^2; P = 0.008 \), with values for females, on average, being 3-1 g/l higher than for males (\( P<0.001 \)). Albumin showed no significant relationship with age or sex. Regression analysis showed that IgG increased by 0.26 g/l per month (\( P<0.001 \)) and the change was consistent in both males and females. AGP showed a significant quadratic effect with age in both sexes, increasing up to about 10 months of age followed by a slight downward trend in older infants (\( y = 620 + 0.043 \times \text{age} + (-0.001) \times \text{age}^2; P<0.001 \)). GSIgM titres showed a significant, but small, downward trend with age (\( B = -0.023; P<0.001 \)). Females had higher titres on average (\( B = 0.134; P = 0.011 \)) but the pattern was not consistent over the total age range (significant interaction effect; \( P=0.004 \)) and males’ GSIgM titres had a significant stronger correlation with age than females (\( r = 0.204 \) and 0.044, respectively, \( \text{t}_{279} = 2.799; P<0.01 \)). The logged L:M ratio also showed a significant downward trend (\( B = -0.006; P=0.003 \)) with, on average, females having a higher L:M ratio (\( B = 0.050; P=0.004 \)).

The relationships between *z*-scores and biochemical variables were examined after controlling for age and sex effects (Table 2). Hb showed positive relationships with all *z*-scores (\( P<0.001, <0.001 \) and 0.004 for HAZ, WAZ and WHZ, respectively); however, albumin showed only a positive relationship with HAZ (\( P=0.016 \) but not with WAZ and WHZ. IgG failed to show any relationships with anthropometric status. The acute-phase protein AGP showed negative relationships with HAZ and WAZ (\( P=0.011 \) and 0.005, respectively) but not with WHZ. *Giardia* infection indicated by the elevation of GSIgM showed significant relationships with WAZ and WHZ (\( P=0.015 \) and 0.039, respectively). Poor intestinal mucosal function (i.e. high L:M ratio) showed significant relationships with lower HAZ and WAZ scores (\( P=0.039 \) and 0.019, respectively).

The inter-relationships between biochemical variables are summarised in Table 3. In general, markers of immune response of inflammation (i.e. IgG, AGP and GSIgM) tended to show significant strong positive relationships. Higher concentrations of albumin were also associated with higher concentrations of the inflammation markers. The L:M ratio, however, did not show significant association with *Giardia* infection or any of the immune responses.

Discussion

Childhood malnutrition remains an almost invariable characteristic of many developing countries. The causes are associated with the complex interplay between poverty, inadequate food quantity and quality, frequent infections, poor sanitary conditions, inappropriate hygiene practices and poor accessibility and affordability of health services. In addition, the capacity to digest and absorb food may be impaired by frequent and chronic illness such as protozoa and geohelminth infection in the intestine. The present study focused on the critical first 2 years of life and specifically investigated the relationship between the infants’ growth status and the level of small-intestinal mucosal damage, parasite burden and the local and systemic inflammation and the immune responses.

The present study found a sharp decline in all three *z*-scores from about 3 months of age. These worsening *z*-scores over the first year of life are in agreement with three other Bangladesh studies, two conducted in urban slums by Arifeen et al. (26) and Karim & Mascie-Taylor (27) and one in a rural area by Begum (28), which showed improvements up to age 4–5 months followed by marked declines, particularly for HAZ and WAZ, until 12 months of age. This downward trend and the timing of failing growth status in Bangladesh are also following, in general, the findings of declines in *z*-scores in developing countries in Africa, Latin America and Asia by Shrimpton et al. (2).

IgG levels generally increase with age in childhood (29), especially in developing countries, and the rise is even faster as production of IgG in the body is stimulated by infections. Fig. 2 compares the relationship between IgG and age in a UK reference (30), a Gambia study (Campbell et al. (31); PG Lunn, personal communication) and the present study. The UK reference and the present study show remarkably similar parallel regression lines (\( B = 0.244 \) and 0.230, respectively) but different intercept values, with a higher level of about 3 units in the present study. The relationship with age in The Gambia was significantly steeper (\( B = 0.581 \)) than the UK or the present study (\( P=0.001 \) and 0.007, respectively), indicating that the Gambian infants were exposed to more
antigens, such as toxins and organisms, and responded by a rise in IgG levels. In the Gambian infants, IgG levels were negatively associated with age-corrected growth rates ($r = -0.63; \ P < 0.001$) (5). However, the present study did not show significant relationships between the IgG level and infants' growth, although all were negative (Table 2).

It is generally assumed that the concentration of AGP at birth is very low, but by 10 months of age (31) infants are thought to have reached adult levels (range 0.5–1.0 g/l (30); mean 0.88 (SD 0.21) g/l (32)). The present study also showed evidence of increasing AGP levels in the first year of life (Fig. 1). Several different assay techniques have been used and there is no agreed cut-off signifying an elevated level for AGP, for example, cut-offs of above 0.75 g/l (33), 1.0 g/l (34) and 1.2 g/l (35–37). In the present study, 60% of infants had an AGP level above 0.75 g/l, 30% were above 1.0 g/l and 16% were above 1.2 g/l and there were increased levels of stunting and underweight in infants with higher concentrations of AGP. This finding is in accord with Filteau et al. (38) who also reported that a higher plasma concentration of AGP was significantly negatively correlated with HAZ as well as WAZ in Ghanaian children below 5 years of age.

Intestinal permeability values in the Bangladeshi infants were higher than those found in a group of UK infants (geometric mean 0.12) (39), and nearly three-quarters of them

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**Fig. 1.** Relationships of z-scores and biochemical variables with age in male (●) and female (○) infants. (a) Height-for-age z-score (HAZ), (b) weight-for-age z-score (WAZ), (c) weight-for-height z-score (WHZ), (d) Hb, (e) albumin, (f) IgG, (g) α-1-acid glycoprotein (AGP), (h) Giardia-specific IgM (GSIgM), (i) lactulose: mannitol (L:M) ratio. Values are means, with 95% CI represented by vertical bars.
(62%) showed evidence of intestinal mucosal damage compared with the UK level. The geometric mean of the L:M ratio in the infants was 0·15 (Table 1), which is lower than the 0·23 found in a previous rural Bangladesh study(40) although the children were older (2–5 years of age) and much lower than the Gambian mean of 0·38 (39). Intestinal mucosal malfunction was also significantly associated with chronic malnutrition (HAZ and WHZ; P<0·039 and 0·019, respectively) rather than acute malnutrition. A previous study in Bangladesh(50) with older children aged 2–5 years also found that better small-intestinal mucosal function was associated with improved WAZ after 8 and 12 months (r=0·19, P<0·002 and r=0·21, P<0·05 for the changes at 8 to 12 months, respectively).

Giardiasis causes acute diarrhoea, so it is not surprising that Giardia infection was related to both acute weight loss in the present study (i.e. WAZ and WHZ; P<0·015 and 0·039, respectively); Lunn et al. in The Gambia found that a short-term weight reduction was associated with elevation of GS1gM(41). However, the present study did not show a clear impact of high endemicity of Giardia infection on intestinal mucosal damage as measured by the L:M ratio.

In the present study, the prevalence of geohelminths in these infants ascertained by eggs in stools was consistently very low (2.8% for Ascaris, 0.8% for Trichuris and no infection of hookworms) throughout the study. The egg counts of Ascaris and Trichuris infection were also very low according to the WHO classification(42), even taking into account the posited lowered fecundity of female Ascaris worms in Bangladesh(43). It is therefore unlikely these very low levels of geohelminth infection would be important to the growth faltering observed in these infants.

The prevalence of Giardia using the GS1gM titre was consistently high, at 98%, in the study. The very high endemicity suggested by the Giardia antibody titre contrasted greatly with the prevalence of Giardia cysts detected in stool examinations, which was very low during the study period. It is generally accepted that excretion of cysts is intermittent; thus a single stool sample, as used in the study, from an infant with a low-intensity infection might contain few, if any, cysts, leading to under-diagnosis(44). It is also important to appreciate that the two techniques are measuring different things: the GS1gM titre is a measure of the body’s exposure to Giardia antigen whereas cysts are only produced when there is an active infection of Giardia trophozoites established in the intestine. These apparently contradictory results may be due to an interaction with breast-feeding, since anti-Giardial factors in the milk may inhibit trophozoite adherence to enterocytes, so reducing the likelihood of persistent infection(45), whilst milk-borne leucocytes actively kill the organisms(46). Consequently, breast-feeding children may be frequently exposed to this parasite but rarely develop a full episode of Giardiasis, so accounting for the differences between stool and serological estimates of Giardia prevalence.

In conclusion, infants in the present study showed a sharp decline in growth and nutritional status in the first 2 years of life and their immune system was frequently stimulated by infections such as Giardia. Infants also had damaged intestinal mucosal function and Giardia infection, in particular, and damaged mucosal function both associated with acute and chronic malnutrition. However, intestinal permeability levels were not directly associated with any of the inflammation markers or with Giardia infection. The elevation of Giardia-specific titres throughout the study is indicative of

### Table 2. Relationships between z-scores and biochemical status after controlling for age and sex

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</table>

Hazard, height-for-age z-score; WAZ, weight-for-age z-score; WHZ, weight-for-height z-score; AGP, a-1-acid glycoprotein; GS1gM, Giardia-specific IgM titre; L:M, lactulose:mannitol.

* B is the regression coefficient. Hb and AGP showed significant quadratic age effects, therefore quadratic age factor also controlled with their linear age factors.
endemicity of *Giardia* infection. So, rural Bangladeshi infants are being exposed to high levels of infection with concomitant gut damage and growth faltering, presumably arising from inappropriate infant-feeding practices and poor standards of hygiene; a previous study in Bangladesh reported that less than 5% of infants below 6 months of age were exclusively breast-fed\(^{(47)}\). Bangladeshi mothers need to be much more aware of the importance of exclusive breastfeeding for the first 6 months as well as the necessity of better feeding and hygiene standards.

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