Sugar malabsorption in healthy neonates estimated by breath hydrogen

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SUMMARY Carbohydrate malabsorption in 110 healthy, term neonates was studied by estimating expired hydrogen \((H_2)\) before and after a feed on day 6 or 7. Carbohydrate malabsorption was assumed to be present if the infant excreted >20 parts per million (ppm) \(H_2\). The frequency of carbohydrate malabsorption in 49 breast-fed infants was 25% \((n=12)\), in 35 infants fed a 7.5% lactose formula it was 31% \((n=11)\), in 26 infants fed a formula containing 1% lactose and 7.3% maltodextrin it was 15% \((n=4)\). These differences in frequency were not significant. Peak \(H_2\) concentration of the malabsorbers in each group, indicating the degree of carbohydrate malabsorption, was 64, 52, and 32 ppm respectively. The degree of carbohydrate malabsorption did not differ significantly between the breast-fed and the high lactose formula groups, but both groups differed from the low lactose group. \(H_2\) excretion was studied for 5 months in an exclusively breast-fed infant. In the first 2 months high concentrations were found and the infant produced 3–5 stools a day. In the next 3 months however, most \(H_2\) estimations were normal and only 1–2 stools a week were passed. With the introduction of solids, daily bowel movements promptly reoccurred. Frequency of carbohydrate malabsorption in newborn infants is fairly high and is primarily related to the lactose intake. The frequency and degree of carbohydrate malabsorption were comparable in breast-fed infants and in infants fed on a high lactose formula; this differs from results previously reported.

The occurrence of carbohydrate malabsorption (CHM) in normal, term neonates on breast milk or cows’ milk-based formulas, may be assumed from the finding of more than 0.5% reducing substances in the faeces, or from the finding of more than 0.15 g/l reducing substances in the urine. The mean ‘normal’ excretion of lactose in the urine of 88 neonates between the 3rd and 7th day of life ranges between 0.2 and 0.5 g/l. Sugar chromatography on stools with excess reducing substances generally shows the presence of lactose, glucose, and galactose. This pattern is consistent with incomplete lactose absorption as nonabsorbed lactose is split by the faecal flora into its component monosaccharides. Lactose malabsorption has been reported to be more common in breast-fed than in bottle-fed infants, even if lactose concentration in the milk is similar. These findings may have practical implications from the nutritional point of view and may help to establish a nonpathogenic type of large bowel microflora. However, the reported differences of CHM in bottle- and breast-fed infants may have been affected by differences in bacterial fermentation of the nonabsorbed sugars and by contamination of the stools with lactose-containing urine.

It has been shown that malabsorption of lactose and other sugars can easily be detected by estimating \(H_2\) in expired air in adults or children. Using a procedure for interval sampling of mixed expired air adapted for infants, we have studied the frequency and the degree of CHM in 110 newborn infants, fed on one of three feeding regimens. The main objective was to find out whether the reported difference in CHM between bottle-fed and breast-fed infants could be confirmed.

Patients and methods

We examined 110 healthy, term infants in the maternity ward of our hospital. Informed consent was obtained from all the parents. Each infant was fed exclusively one of three milks: breast milk \((n=49)\), cows’ milk-based formula containing 7.5% lactose \((\text{Almiron M2-B*}) (n=35)\), or 1% lactose and 7.3% maltodextrin \((\text{Almiron A-B*}) (n=26)\). The composition of each formula is shown.

*Manufactured by Nutricia, Zoetermeer, the Netherlands.
in Table 1. Infants on mixed feeds were not studied, but breast-fed infants whose feeding had been supplemented with glucose solution during the first few days after birth were included. Mothers who breast fed their babies were instructed to avoid fruit and fruit juices on the day before and during the examination since fruit sugars may be excreted into the breast milk.

To avoid the influence of large differences in intakes of milk/kg, the actual body weights and ingested volumes of the feeds were recorded. In the breast-fed group ingested volume was estimated by test weighing the infant before and after the feed. Serial H₂ estimations were done on day 6 or 7, starting at 0900 hours, three hours after the first feed that day. The second breath sample was taken at 1000 hours (just before the second feed), the third at 1100 hours, and the fourth at noon. Duplicate breath samples were stored in Vacutainers for gas chromatographical analysis. Details of the procedure have been presented previously.¹⁶ In children with lactose malabsorption, an increase above the baseline value of >10 ppm H₂ correlated positively with the occurrence of clinical symptoms.¹⁶-¹⁷ Since the frequent feedings in the study group did not allow H₂ excretion to be measured during fasting, and this is below 10 ppm in older children (R G Barr, 1979, personal communication), a total concentration of 20 ppm H₂ in mixed expired air was assumed to be a safe indication for the presence of CHM in these neonates.

Results

All 27 infants with CHM were white, 7 (26%) of them weighed more than 3500 g, 13 (48%) weighed more than 3000 g, and 7 (26%) weighed more than 2500 g, the lowest weight being 2575 g. No infant had diarrhoea although all infants passed at least two loose stools a day. The mean volume of milk ingested (ml/kg) did not differ between the three feeding groups and only small individual variations were found.

Most of the infants with CHM had raised H₂ concentrations in the first breath sample. This was generally followed by a further increase, although in some infants the concentration remained more or less constant, and 2 infants had decreasing concentrations towards the end of the sampling period. Peak H₂ excretion for each infant is recorded in Fig. 1. The frequency of CHM in the 49 breast-fed infants was 25% (n=12), 31% in the 35 infants fed 7.5% lactose formula (n=11), and 15% in the 26 infants fed 1% lactose formula with 7.3% maltodextrin (n=4). These frequencies are not statistically different. The relative degree of CHM, measured as peak H₂ excretion, was no different for breast-fed infants than for those on the 7.5% lactose formula. But both of these groups had a significantly higher degree of CHM compared with the group on 1% lactose and 7.3% maltodextrin formula (P<0.01, Table 2).

H₂ excretion in the breast-fed infant during the
first 150 days of life changed during the day (Fig. 2), and varied greatly on different days (Fig. 3). The highest concentrations however, were found in the first 2 months. Stool frequency in the first 2 months was 3–5 a day. In the next 3 months, while the baby was still being exclusively breast fed, H₂ excretion became normal or was moderately raised (Fig. 3). Only 1–2 stools a week were passed in the last 3 months and no sign of constipation occurred. Introduction of solids at age 6 months immediately resulted in daily passage of 1–2 well-formed stools.

Table 2  Frequency and relative degree of carbohydrate malabsorption in three groups of neonates on different feeding regimens

<table>
<thead>
<tr>
<th></th>
<th>Breast milk (n=49)</th>
<th>Almiron M2-B (n=35)</th>
<th>Almiron A-B (n=26)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of infants (%)</td>
<td>12 (25)</td>
<td>11 (31)</td>
<td>4 (15)</td>
</tr>
<tr>
<td>95% confidence limits for percentage malabsorbers</td>
<td>15–39</td>
<td>18–49</td>
<td>7–34</td>
</tr>
<tr>
<td>Mean ppm H₂ malabsorbers</td>
<td>64</td>
<td>52</td>
<td>32 (P&lt;0.01)</td>
</tr>
<tr>
<td>SD</td>
<td>43</td>
<td>23</td>
<td>9</td>
</tr>
</tbody>
</table>

Horizontal bar gives upper normal value.

Fig. 3  H₂ excretion over 5 months in an exclusively breast-fed infant.

Discussion

This study confirms earlier observations, based on the finding of excess reducing substances in the stools (>0.5 %) or in the urine (>0.15 g/l), that many neonates have some degree of CHM. The amount of sugars found in excreta gives some indication of the degree of CHM; this tended to be higher in breast-fed infants than in bottle-fed ones. Conclusions about the different degree of CHM between these groups have not been made.

It has been shown previously that H₂ production is primarily related to the amount of fermentable substrate in the colon and this amount can be calculated from the expired H₂. H₂ response to the same dose of lactose may differ considerably between individual lactose malabsorbers, yet, the mean H₂ response for a group of lactose malabsorbers to different doses of lactose differs significantly. It therefore seemed justified to use mean H₂ concentrations for different feeding groups to find out the effect of diet on the degree of CHM. Unlike reports based on an estimation of faecal sugar, the degree of CHM as shown by mean H₂ response did not differ between breast-fed infants and those on the 7.5% lactose formula, but CHM in both these groups was different from that in the group on the 1% lactose 7.3% maltodextrin formula (P<0.01).

Reports concerning the frequency of CHM in neonates are conflicting, and figures range from 0 to 50%. The precise frequency of CHM in neonates is difficult to establish as CHM may be
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present intermittently, as judged from the lactose content in urine and the excretion of H₂ (Figs 2 and 3), and data from former studies as well as from this one are cross-sectional (Fig. 1). The actual frequency may therefore be considerably greater than these studies suggest. This suggestion is supported by the findings of our longitudinal study (Fig. 3). A relationship may exist between CHM and frequency of defecation; this is suggested by the pronounced decrease in stool frequency, synchronous with the decreasing concentrations of expired H₂ after age 2 months (Fig. 3).

Lactose is often implicated as the major source of excess faecal and urinary sugar in newborn infants although other sugars have also been found. The accumulating evidence for lactose malabsorption in a considerable proportion of newborn infants seems to contradict the finding that brush border lactase activity is fully mature by the time of birth although its maximum is reached only shortly before delivery. Presumably the total splitting capacity of the neonatal small bowel is reduced because in these infants the surface area is smaller than in older infants. Previous studies that have been based on an estimation of faecal sugar have stressed the influence of the nature of the feeding, and have suggested that frequency and degree of CHM in breast-fed infants were higher than in formula-fed neonates, even when these formulas contained lactose in amounts comparable with human milk. Our study failed to confirm this interpretation since frequency of abnormal H₂ excretion in breast-fed infants did not differ from that in bottle-fed infants, and was even lower in the breast-fed group compared with the high lactose formula group (Table 2).

Our findings indicate that CHM in healthy, term neonates is primarily due to an age-related maturation process of lactose digestion and absorption, rather than to the nature of the feeding.

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References


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