Amylin peptide levels are raised in infants of diabetic mothers

V Kairamkonda, A Deorukhkar, R Coombs, R Fraser, T Mayer

ORIGINAL ARTICLE

Background: Amylin is a novel 37 amino acid peptide hormone that is co-secreted with insulin from the pancreas in response to food intake. As a potent inhibitor of gastric emptying it plays an important role in the control of carbohydrate absorption. Feed intolerance is common in infants of diabetic mothers (IDM). Aims: To establish a normal range of amylin levels in healthy neonates, and to determine whether serum amylin levels are raised in IDM.

Methods: A serial sample of 221 infants >28 weeks gestation was enrolled prior to delivery over a 12 month period. Blood samples collected immediately after birth (umbilical cord), and at the routine Guthrie test were analysed for amylin and insulin levels.

Results: Amylin levels in umbilical cord (n=181) and Guthrie samples (n=33) of healthy infants were 5.7 (3.0–9.1) and 6.9 (2.9–9.0) pmol/l respectively. IDM had significantly raised amylin levels in both cord (n=31; 32.7 pmol/l, 25.9–48.1) and Guthrie samples (n=8; 18.1 pmol/l, 15.3–23.6). Amylin correlated positively with insulin (n=42; r=0.67; 95% CI 0.4 to 0.81), birth weight (r=0.22; 95% CI 0.08 to 0.36), and gestation (r=0.18; 95% CI 0.03 to 0.32). Umbilical cord venous amylin levels showed agreement with arterial cord amylin levels (n=34, mean bias −0.2, 95% CI 3.1 to −3.6).

Conclusions: Amylin levels are significantly increased in the umbilical cord and Guthrie blood samples in IDM.

SUBJECTS AND METHODS

A serial sample of 221 infants was enrolled prior to delivery at a tertiary maternity hospital from March 2003 to February 2004, over a 12 month period. This study was approved by the South Sheffield Research Ethics Committee and required formal written parental consent. Inclusion criteria were healthy infants born at a gestational age >28 weeks and infants born to diabetic mothers (including gestational, insulin dependent, and non-insulin dependent diabetes mellitus). Infants with congenital anomalies and deliveries associated with chorioamnionitis or significant antenatal or labour complications were excluded. Preterm infants, ventilated for poor respiratory effort, who were clinically stable, were not excluded.

Prospective serial umbilical cord blood samples were collected immediately after birth, and at the routine Guthrie test (postnatal day 5). Guthrie samples from many of our planned recruits in the community were not feasible.
due to strict collection and temperature requirements for blood collection. Paired umbilical cord, venous, and arterial samples were obtained whenever possible from inpatient infants. All samples were analysed for amylin, and where volume of blood was sufficient, insulin assay was carried out to confirm co-secretion. A 1 ml sample of blood was drawn into a tube containing EDTA with 500 IU of aprotinin; it was kept on ice before separation and storage of plasma at −70°C. Plasma total amylin immunoreactivity was quantified using a monoclonal antibody-based sandwich immunoassay system developed by LINCO Research, Inc. (Missouri, USA).

This has sensitivity of 1 pmol (50 μl plasma sample size). Inter- and intra-assay coefficients of variation at 50 pmol spiked concentration of amylin were 13–18% and 2.5–5% respectively. Plasma immunoreactive insulin was measured by enzyme-linked immunosorbent assay that has inter- and intra-assay coefficients of variation of 5.6% and 5.3% (BioSource Labs, Belgium).

Relevant information was retrieved from an electronic patient database. Infant data included birth weight, gestation, and gender. Maternal data included medical, antenatal, and labour history.

Statistical analysis
Statistical analysis was performed using Medcalc version 7.4.1.2 1993–2004. A priori statistical advice suggested a sample size of at least 200 observations (cord blood samples) for the calculation of a reference interval. To detect a difference in means of 15 and assuming a standard deviation of 13, at least 21 IDM were required (5% two-sided significance level and 90% power). As the data were non-normal, non-parametric tests were employed. Data are presented as median values with interquartile range (25th–75th centile). Pearson’s correlation coefficient was used to assess correlation between amylin and insulin levels. A p value of less than 0.05 was considered significant. A Bland–Altman plot of difference between the methods (y axis) against their average (x axis) was used to measure agreement between the two sampling methods (cord arterial and cord venous) (fig 4). This method aims to investigate a possible relation between measurement error and the true value which is unknown. Therefore the mean, average discrepancy (bias), trend, and scatter of the two measurements are used as a guide to the comparability of the two methods.

RESULTS
A total of 221 infants were enrolled, of which 31 were infants of diabetic mothers. Amylin levels were analysed on venous umbilical cord blood from 181 healthy infants; nine infants were excluded from this group as cord blood obtained was unsuitable for analysis. Additional serum amylin levels were obtained from 33 of the healthy infant group on the fifth postnatal day (Guthrie). Among infants born to diabetic mothers (n = 31), Guthrie blood samples were obtained for amylin in eight infants, who were all inpatients at that time (fig 1).

In the population of healthy neonates the median cord serum amylin level was 5.7 pmol/l (3.0–9.1). A similar value was observed on the fifth postnatal day (table 1).

Male amylin level in cord blood from infants of diabetic mothers was significantly higher (32.7 pmol/l (25.9–48.1)) compared to healthy newborn infants (5.7 pmol/l (3.0–9.1)) (table 2).

In infants of diabetic mothers, the amylin levels were lower on postnatal day 5 (18.1 pmol/l (15.3–23.6)) compared to cord blood, but were still significantly higher than values in healthy neonates (6.9 pmol/l (2.9–9.0)) at the same postnatal age (table 3).

Cord serum amylin levels correlated positively with gestational age and birth weight (figs 2 and 3). Amylin levels from arterial and venous cord blood were comparable when paired samples were obtained for analysis (n = 34, r = 0.98, 95% CI 0.96 to 0.99, p < 0.0001). The Bland–Altman plot (fig 4) shows the mean difference between the sampling methods to be −0.3 pmol/l and the average discrepancy (bias) to be −0.2. Thus the venous amylin levels tend to be lower, but despite this the limits of agreement (−3.6 and 3.1) are small enough for us to be confident that cord arterial samples may be used interchangeably with cord venous samples for measuring amylin levels. Finally, a positive correlation was observed between amylin and insulin levels (n = 42, r = 0.67, 95% CI 0.4 to 0.81, p < 0.0001).

DISCUSSION
Amylin is co-secreted with insulin and is a potent inhibitor of gastric motility. High serum amylin levels have been reported in early type 2 diabetes and critically ill children with feed intolerance. The diurnal profile of serum amylin levels has been established in healthy children and adults. This is the first study to establish the normal range of serum amylin in healthy neonates at birth and at the fifth postnatal day. The serum amylin levels in healthy neonates in our study (5.7 pmol/l) correspond to those observed (mean (SD)) in the paediatric (5.0 (1.94) pmol/l) and adult (5.0 (0.4) pmol/l) populations. It is possible that the cord amylin levels represent transplacental passage of this peptide. Although feasible, this seems unlikely as there was no fall in serum amylin levels by the fifth postnatal day in healthy infants when compared to levels obtained at birth. The high

### Table 1 Amylin levels and demographic data of healthy infants in the cord and Guthrie groups

<table>
<thead>
<tr>
<th></th>
<th>Healthy infants</th>
<th>Cod (n = 181)</th>
<th>Guthrie (n = 33)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amylin (pmol/l)</td>
<td>5.7 (3.0–9.1)</td>
<td>6.7 (2.9–9.0)</td>
<td></td>
</tr>
<tr>
<td>Gestation (wk)</td>
<td>38 (37–39)</td>
<td>35 (34–38)</td>
<td></td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>3.12 (2.58–3.56)</td>
<td>2.06 (1.65–3.13)</td>
<td></td>
</tr>
<tr>
<td>Gender (M:F)</td>
<td>1.1:1 (97:84)</td>
<td>1:1 (17:16)</td>
<td></td>
</tr>
</tbody>
</table>

IDM versus Guthrie (healthy infants): p = 0.0001. Values are presented as medians (25th–75th centile).

### Table 2 Amylin levels and demographic data of IDM and healthy infants in the cord group

<table>
<thead>
<tr>
<th></th>
<th>IDM</th>
<th>Healthy infants</th>
</tr>
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<tbody>
<tr>
<td>Amylin (pmol/l)</td>
<td>32.7 (25.9–48.1)</td>
<td>5.7 (3.0–9.1)</td>
</tr>
<tr>
<td>Gestation (wk)</td>
<td>37 (34.0–39.0)</td>
<td>38 (37.0–39.0)</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>3.31 (2.72–3.84)</td>
<td>3.12 (2.58–3.56)</td>
</tr>
<tr>
<td>Gender (M:F)</td>
<td>1.2:1 (17:14)</td>
<td>1.1:1 (97:84)</td>
</tr>
</tbody>
</table>

IDM versus healthy infants (cord): p = 0.0001. Values are presented as medians (25th–75th centile).

### Table 3 Amylin levels and demographic data of IDM and healthy infants in the Guthrie group

<table>
<thead>
<tr>
<th></th>
<th>Guthrie (n = 8)</th>
<th>Healthy infants (n = 33)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amylin (pmol/l)</td>
<td>18.1 (15.3–23.6)</td>
<td>6.9 (2.9–9.0)</td>
</tr>
<tr>
<td>Gestation (wk)</td>
<td>35 (33–36)</td>
<td>35 (34–38)</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>2.94 (1.97–3.56)</td>
<td>2.06 (1.65–3.13)</td>
</tr>
<tr>
<td>Gender (M:F)</td>
<td>1.6:1 (5:3)</td>
<td>1:1 (17:16)</td>
</tr>
</tbody>
</table>

IDM versus healthy infants (Guthrie): p = 0.0001. Values are presented as medians (25th–75th centile).
Amylin peptide levels in infants of diabetic mothers

Amylin levels in IDM further support the theory that transplacental passage does not occur as serum amylin levels are subnormal in diabetic patients.10,11,14,15 Thus these high levels obtained in IDM would seem to represent endogenous amylin production. Additionally studies performed in perfused human placenta with pramlintide, a synthetic analogue of amylin, indicate that pramlintide has low potential (ratio $t_{1/2} = 50$ minutes), which is primarily metabolised by the kidneys after subcutaneous dosing, would further question transplacental passage being responsible for the observed serum amylin levels. In summary, this evidence would suggest that the serum amylin levels in the neonatal period are due to endogenous production, and are in the range reported in the paediatric and adult populations.

In this study serum amylin levels obtained from cord blood were significantly raised in infants born to diabetic mothers. These levels are comparable to those reported in critically ill children with feed intolerance.16,17 We suggest that these high serum amylin levels observed in IDM may be partially responsible for the feed intolerance seen in this population. This is in keeping with the physiological action of amylin, as it is 15–20 times more potent than other known inhibitors of gastric motility.18 There are a number of possible explanations for such high levels observed in IDM. Amylin is deficient in type 1 diabetes mellitus and is present in excess in conditions in which insulin is hyper-secreted, for example in insulin resistance states.19 It would therefore seem plausible that transient neonatal hyperinsulinaemia in IDM due to in utero exposure to maternal hyperglycaemia, may also stimulate an increase in release of endogenous amylin. This is further supported by evidence of co-secretion of amylin and insulin, shown by the positive correlation between amylin and insulin in our study $(r = 0.67, 95\% \text{ CI } 0.40$ to $0.81, p < 0.0001)$ and previous studies.14 Unfortunately we were unable to obtain sufficient sample volume for analysis of paired amylin and insulin on the fifth postnatal day. Amylin has been shown to have both vasodilator20 and anti-inflammatory21 properties. It shares 50% homology with calcitonin gene related peptide, which is recognised to be raised in acute inflammatory states such as trauma,4 sepsis (neonatal and adult),2 and hypotensive shock.16 We therefore excluded infants born to mothers with chorioamnionitis and those with significant antenatal and labour complications.

In IDM, the fifth postnatal day amylin levels were lower than cord samples at birth. This gradual decline possibly represents absence of the hyperinsulinaemic environment and would coincide with the time course of resolution of the feed intolerance observed in IDM. However, our study did not attempt to correlate amylin levels with objective measures of gastric emptying and feed intolerance or its resolution. This is an important limitation of our study. The observed serum amylin levels in IDM at the fifth postnatal day were still above our proposed normal range for healthy infants. In retrospect, samples over a longer time period may have been useful and should be considered in the design of future studies.

Infants in the Guthrie group were lighter in weight and of a lower gestational age than in those in cord blood group. This is possibly due to a selection bias towards blood collection of inpatient neonates. Agreement was shown between paired cord arterial and venous amylin levels; therefore either of the two methods could be used to measure amylin. This observation will help in future studies involving umbilical blood sampling.

![Figure 2: Correlation between cord amylin and gestation (n=181). $r=0.20$, 95% CI 0.04 to 0.35, $p=0.012$.](image)

![Figure 3: Correlation between cord amylin and birth weight (n=181). $r=0.21$, 95% CI 0.05 to 0.36, $p=0.002$.](image)

![Figure 4: Method comparison (Bland-Altman) of paired venous and arterial levels. n=34, mean bias $-0.2$, limits of agreement $3.1$ to $-3.6$.](image)
We observed a direct correlation between amylin levels with ascending birth weight and gestation. A similar finding has been reported with the calcitonin gene related peptide.7 The significance of this observation is unclear.

The results of this study will be useful for the ongoing study investigating the role of amylin in preterm infants with feed intolerance. Amylin has been shown to be raised in rat intestinal ischaemic injury,22 and may have a biologic role in sepsis.9 Future research in the role of amylin in necrotising enterocolitis and sepsis in the neonatal population may benefit from normal values established by our study. AC187 is a truncated amylin peptide antagonist that acts by selectively and potently blocking amylin receptors.9 The biological actions of amylin in IDM and preterm infants with delayed gastric emptying need to be elucidated prior to evaluation of pharmacological blockade of this peptide as a potential therapeutic option.

In conclusion, we have established a normal range for amylin levels in neonatal population and presented evidence in support of our hypothesis that amylin levels are raised in IDM.

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REFERENCES