Pregnancy glycaemia and cord-blood levels of insulin and leptin in Pakistani and white British mother-offspring pairs

Citation for published version:
Lawlor, DA, West, J, Fairley, L, Nelson, SM, Bhopal, RS, Tuffnell, D, Freeman, DJ, Wright, J, Whitelaw, DC & Sattar, N 2014, 'Pregnancy glycaemia and cord-blood levels of insulin and leptin in Pakistani and white British mother-offspring pairs: findings from a prospective pregnancy cohort' Diabetologia. DOI: 10.1007/s00125-014-3386-6

Digital Object Identifier (DOI):
10.1007/s00125-014-3386-6

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published in:
Diabetologia

Publisher Rights Statement:
This article is distributed under the terms of the Creative Commons Attribution License which permits any use, distribution, and reproduction in any medium, provided the original author(s) and the source are credited.

General rights
Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.
Pregnancy glycaemia and cord-blood levels of insulin and leptin in Pakistani and white British mother–offspring pairs: findings from a prospective pregnancy cohort

Debbie A. Lawlor · Jane West · Lesley Fairley · Scott M. Nelson · Raj S. Bhopal · Derek Tuffnell · Dily J. Freeman · John Wright · Donald C. Whitelaw · Naveed Sattar

Received: 4 June 2014 / Accepted: 29 August 2014
© The Author(s) 2014. This article is published with open access at Springerlink.com

Abstract
Aims/hypothesis To determine the extent to which gestational fasting and postload levels of glucose explain differences in infant fat mass between UK-born Pakistani and white British infants.

Methods Analyses were undertaken in a prospective pregnancy cohort study of 1,415 women and their singleton live-born infants (629 white British and 786 Pakistani). Infant fat mass was assessed by cord-blood leptin levels and fetal insulin secretion by cord-blood insulin levels. Maternal OGTTs were completed at 26–28 weeks of gestation.

Results Pakistani women had higher fasting and postload glucose levels and greater incidence of gestational diabetes than white British women. Higher fasting and postload glucose levels were associated with higher cord-blood insulin and leptin levels in all participants, irrespective of ethnicity. Cord-blood leptin levels were 16% (95% CI 6, 26) higher in Pakistani than in white British infants. After adjustment for fasting glucose levels, this difference attenuated to 7% (−3, 16), and with additional adjustment for cord-blood insulin levels it attenuated further to 5% (−4, 14). Path analyses supported the hypothesis that fasting glucose levels mediate the relationship of Pakistani ethnicity to greater fat mass at birth, as measured by cord-blood leptin levels; on average, 19% of this mediation involvement fetal insulin secretion. Postload glucose levels did not act as an important mediator of ethnic differences in cord-blood insulin levels. Results were very similar when 130 women with gestational diabetes were removed.

Conclusions/interpretation These novel findings suggest a role of maternal pregnancy glycaemia in mediating differences in fat mass between Pakistani and white British infants.

Keywords Birthweight · Cord-blood insulin · Cord-blood leptin · Epidemiology · Ethnicity · Gestational diabetes · Glucose

Abbreviations
BiB Born in Bradford
GDM Gestational diabetes mellitus
Introduction

For a given BMI, South Asian compared with white European adults have greater fat mass, leading to the suggestion that they have a ‘thin-fat’ insulin-resistant phenotype that underlies their increased type 2 diabetes risk [1–3]. Recent evidence suggests that this phenotype is present in children [4–6], and possibly at birth [7]. For a given birthweight, South Asian babies born in India have been found to be fatter (as measured by subscapular and triceps skinfolds or cord-blood leptin levels) than white European babies born in Europe or the USA [7–9]. These differences may reflect different socioeconomic, lifestyle and healthcare characteristics between people living in India compared with those in Western countries. However, we have recently shown similar results when comparing babies of Pakistani origin conceived and born in the UK with white British babies born in the same maternity unit [10].

Greater gestational hyperglycaemia amongst South Asian women [11–14] is a plausible exposure that could contribute to greater fat mass at birth in South Asian infants because (1) greater circulating pregnancy fasting and postglucose concentrations promote fetal insulin secretion and fat accretion in a linear dose–response fashion [15–19]; and (2) an increased risk of gestational diabetes mellitus (GDM), hyperglycaemia and type 2 diabetes in people of South Asian origin is well established [1, 11–14, 20–25].

The aim of this study was to determine the extent to which differences between Pakistani and white British women in gestational glucose levels mediate differences in cord-blood leptin levels (a valid proxy for birth fat mass that has been used in previous studies for this purpose [7, 10, 26, 27]). We also investigated the role of cord-blood insulin levels in these relationships. This work extends our earlier work [10], which did not explore possible mediators of ethnic differences in cord-blood leptin levels at birth. To the best of our knowledge, the role of maternal gestational glycaemia in mediating differences in infant fatness between Pakistani (or any other South Asian group) and white British infants has not been previously explored.

Methods

This study was approved by the Bradford Research Ethics Committee (ref 06/Q1202/48). All participants provided written informed consent. The study was undertaken in a planned pseudo-randomly selected subgroup of the Born in Bradford (BiB) cohort in whom relevant cord-blood outcomes were assayed. BiB is a prospective pregnancy cohort study that collected detailed information from 12,453 women who experienced 13,776 pregnancies resulting in 13,818 births [28]. To be eligible for BiB, women had to be booked to give birth between March 2007 and December 2010 in the maternity department at Bradford Royal Infirmary, in the North of England [28].

BiB women were recruited primarily at their OGTT appointment (26–28 weeks’ gestation). All women booked for delivery in Bradford are offered a 75 g OGTT (comprising fasting and 120 min postload samples) at around 26–28 weeks’ gestation. At recruitment, BiB participants completed an interviewer-administered questionnaire. Interviews were conducted in English or in a range of South Asian languages, including Urdu, Mirpuri, Bengali and Punjabi [28].

To be eligible for the cord-blood substudy, BiB participants had to have a live birth from a singleton pregnancy that occurred between mid-October 2008 and mid-October 2009. Electronic supplementary material (ESM) Table 1 shows that the distributions of participant characteristics are similar in the subgroup used for this study and those in the whole cohort of singleton live births. Figure 1 shows further exclusions from this study. Participants had to be of Pakistani or white British origin. The BiB cohort includes all eligible women, irrespective of their ethnic origin, but for this study other ethnic groups were either too small to be examined separately or were not relevant to the comparison of South Asian and white European ethnicity. Participants who had not completed an OGTT were excluded. Over 80% of the whole BiB cohort who were eligible for an OGTT completed it and had valid data on fasting and postload glucose levels; their characteristics were similar to those of participants who did not complete the OGTT [28]. Any women with pre-existing diabetes were not invited to have an OGTT, thus by excluding women without OGTT results we were also excluding any women with diagnosed existing diabetes. Following these exclusions, there were 1,489 eligible participants for this study and, of these, 1,415 (95%) had complete data on all covariables included in

![Fig. 1 Flow diagram of participants](chart)
any model; these 1415 form the main analysis sample for this study (Fig. 1).

To be defined as Pakistani, women had to state that they belonged to category 10 of the UK Office of National Statistics criteria (Asian/British Asian subcategory and labelled Pakistani) [29]. To be defined as white British, women had to state that they belonged to category 1 (white subcategory and labelled English/Welsh/Scottish/Northern Irish/British). Full details of all of the Office of National Statistics categories are provided in the ESM Methods. For Pakistani women, additional information on their country of birth and that of their partner and their parents was also collected. This confirmed South Asian (Indian, Pakistani or Bangladeshi) familial origin for all who had reported that they were Pakistani. In a previous publication we have shown that differences in cord-blood leptin levels between Pakistani and white British infants were the same irrespective of place of birth of parents and grandparents (i.e. generation of migration of the Pakistani women) [10], and have not explored that further in this paper.

Participants completed a morning OGTT after fasting overnight. Fasting blood samples were taken for immediate analysis. Participants then consumed a standard solution containing the equivalent of 75 g anhydrous glucose over 5 min. After 120 min a second plasma sample was taken. All glucose assays were completed in the same laboratory in Bradford, using a glucose oxidase method, with laboratory staff blinded to the participants’ ethnicity or other characteristics. GDM was defined according to modified WHO criteria operating at the time these women were pregnant [30], as either fasting glucose ≥6.1 mmol/l or 2 h glucose ≥7.8 mmol/l. The modification that was applied in Bradford was using a fasting glucose threshold of ≥6.1 mmol/l. Although at the time the threshold recommended by WHO was ≥7.0 mmol/l, GDM was defined, including by the WHO, as any impaired glucose tolerance diagnosed for the first time in pregnancy, hence we used the threshold indicative of impaired fasting glucose. As a consequence, the definition being used in these women is similar to the recently proposed International Association of Diabetes and Pregnancy Study Group thresholds, which are included in the new WHO guidance [31].

Venous cord-blood samples were obtained at delivery by the attending midwife, following research protocols. Samples were refrigerated at 4°C in EDTA tubes until collected by laboratory staff within 12 h. Samples were then spun, frozen and stored at −80°C. Insulin is known to be stable in EDTA samples at room temperature for over 24 h [32]. The samples were transferred to Glasgow for completion of leptin and insulin assays. Leptin was measured by a highly sensitive in-house ELISA validated against the commercially available Linco assay, as detailed previously [33]. Insulin was measured by an ultrasensitive solid-phase two-site immunoassay ELISA (Mercodia, Uppsala, Sweden) that does not cross-react with proinsulin. Laboratory staff were blinded to the participants’ ethnicity and other characteristics.

Maternal age, education and smoking data were obtained from the interviewer-administered questionnaire completed at recruitment. Maternal education was equivalised to UK standard attainments and participants were included in one of five mutually exclusive categories (full details in the ESM Methods). Maternal smoking was categorised as never, past (but not during this index pregnancy), current/during the index pregnancy. Maternal BMI was calculated from height measured at the time of recruitment and from weight measured at first antenatal clinic, which was abstracted from the antenatal medical records; median (interquartile range) gestational age of the mothers at this booking clinic was 12 (11–14) weeks. Maternal parity, gestational age (to the last completed week) and infant birthweight and sex were abstracted from the antenatal/obstetric medical records.

Statistical analyses Maternal glucose levels and cord-blood levels of insulin and leptin were positively skewed and were described using medians and interquartile range. In regression analyses that used these variables as exposures or mediators (i.e. independent variables) they were used in their original form (mmol/l for glucose and pmol/l for cord-blood insulin) and in those that used them as outcomes they were natural log transformed so that the residuals in all models were approximately normally distributed. The resultant regression coefficients were transformed so that they represent differences in means as a percentage (%) per unit of exposure.

We tested our hypotheses in two ways. First, by a series of separate linear regression models that examined the associations of (1) ethnicity with fasting and postload glucose levels; (2) gestational fasting and postload glucose levels with cord-blood levels of insulin and leptin; (3) cord-blood insulin levels with cord-blood leptin levels; and (4) ethnicity with cord-blood levels of insulin and leptin. In these regression models we adjusted for maternal age, parity, smoking, education and BMI, gestational age and offspring sex. Details of the rationale for adjusting for these covariables are provided in the ESM Methods. In regression analyses for the association between ethnicity and cord-blood leptin levels, we additionally adjusted for potential mediation by fasting/postload glucose levels and cord-blood insulin levels.

Second, we used path analysis to simultaneously run all of the equations from the different regression analyses and pictorially show the standardised regression coefficients between exposures, potential mediators and outcomes. We included all of the same covariables in this path analysis as in the serial regression analysis, although on the figures we show only the results for the main exposure, outcome and mediating pathways. The path analyses were done with all variables standardised to have mean 0 and standard deviation 1 and run using the sem command in Stata.
We examined whether associations of glucose with cord-blood levels of insulin and leptin were linear in each ethnic group using fractional polynomials [34]. Differences between white British and Pakistani participants in the associations between gestational fasting and postload glucose levels with outcomes were tested using an interaction (ethnicity*glucose) test.

The main association analyses were completed in all 1,415 mother–offspring pairs and also in the 1,285 pairs after removal of the 130 pairs in whom the mother met the criteria for GDM.

Results

ESM Table 2 shows characteristics for all participants and separately for Pakistani and white British women. Pakistani women were on average older, had greater parity, were much less likely to smoke and were more likely to be in the lowest and highest educational attainment categories. They had higher gestational fasting and postload glucose levels and lower BMI. Pakistani infants had higher cord-blood levels of insulin and leptin and lower birthweight than white British infants. None of the associations examined in this study differed by offspring sex (all $p_{interaction}$ ≥ 0.2) and all regression analyses are presented with both sexes combined.

Cord-blood leptin level was positively associated with birthweight in both ethnic groups (ESM Fig. 1). For a 1 unit increase in logged cord-blood leptin level, birthweight increased by 311 g (95% CI 267, 355) in white British women and 263 g (95% CI 227, 302) in Pakistani women ($p_{interaction}$ = 0.11). $R^2$ values indicated that cord-blood leptin levels explained 25% and 22% of the variation in birthweight in white British and Pakistani infants, respectively.

One hundred and thirty (9%) of the women were identified as having GDM; it was more common in Pakistani ($n=97$ [12%]) than white British ($n=33$ [5%]) women ($p<0.001$). For the vast majority of women who met the criteria, GDM was diagnosed because of a high postload glucose level, with 77% of Pakistani and 88% of white British women with GDM having a postload glucose level of ≥ 7.8 mmol/l but a fasting glucose level in the normal range.

Figure 2 shows the scatter plots and unadjusted predicted regression lines for the associations of fasting glucose with levels of cord-blood insulin (Fig. 2a) and cord-blood leptin (Fig. 2b) in all participants. ESM Fig. 2 shows similar results for postload glucose and ESM Fig. 3 indicates the results when women with GDM are removed. In both Pakistani and white British participants, and irrespective of whether women who met criteria for GDM were removed or not, there were linear increases in levels of cord-blood leptin and cord-blood insulin across the distribution of increasing maternal gestational levels of fasting and postload glucose. The fractional polynomial models supported linear associations of both fasting and postload glucose levels with cord-blood levels of insulin and leptin in both Pakistani and white British women (all $p$ values ≤ 0.001 for linear association and >0.2 for any power terms).

The linear positive associations of maternal gestational levels of fasting and postload glucose with cord-blood levels of insulin and leptin remained with adjustment for confounders in both the whole cohort (ESM Table 3) and in those who did not meet criteria for GDM (ESM Table 4). For cord-blood levels of insulin and leptin, the magnitudes of association were greater with fasting than with postload glucose levels (Table 1). Associations of fasting glucose levels with cord-blood levels of leptin and insulin were somewhat stronger when women with GDM were removed from the analyses than when all participants were included in analyses (Table 1). After adjustment for confounding factors, the magnitudes of association of fasting and postload glucose levels with cord-blood levels of insulin and leptin were similar in the two ethnic groups. Cord-blood insulin level was positively associated with cord-blood leptin level in both ethnic groups, again with similar magnitudes of association between the two ethnic groups (Table 1).
Higher cord-blood insulin levels in Pakistan compared with white British infants remained after adjustment for confounders in both the whole cohort and when women with GDM were removed (Table 2). Adjustment for potential mediation by maternal fasting glucose levels attenuated the confounder-adjusted difference completely to the null in the whole cohort and markedly when those with GDM were excluded, whereas adjustment for postload glucose levels reduced the difference only slightly.

Cord-blood leptin levels remained higher in Pakistani than white British infants after adjustment for confounders, being 16% (95% CI 6, 26) and 12% (95% CI 4, 21) higher in the whole cohort and the cohort without GDM, respectively. In both cohorts, adjustment for potential mediation by fasting glucose levels reduced the ethnic difference by approximately 50% and statistically the difference was consistent with the null, whereas adjustment for postload glucose levels resulted in very little mediation. With additional adjustment (for confounders and mediation by fasting glucose levels and cord-blood insulin levels), there was a further attenuation to the null of the ethnic difference in cord-blood leptin levels (Table 2).

Figure 3 shows the results of the path analyses for the ethnic difference in cord-blood leptin levels in the whole cohort (with equivalent results when those with GDM were removed shown in ESM Fig. 4). The confounder-adjusted partial correlation coefficient for the difference in cord-blood leptin levels between Pakistani and white British women was 0.058 (p<0.001), i.e. Pakistani infants had levels of cord-blood leptin that were 0.058 of an SD higher than white British infants. This difference is calculated from the path analysis (Fig. 3) by adding together the two paths from ethnicity to cord-blood leptin level: one that goes from ethnicity through fasting glucose level to cord-blood leptin level (0.18×0.26=0.047) and one that goes from ethnicity to fasting glucose level, and from fasting glucose level to cord-blood insulin level and from there to cord-blood leptin level (0.18×0.19×0.31=0.011). The path analysis shows that the difference in cord-blood leptin level between Pakistani and white British infants was completely explained by higher maternal fasting glucose levels amongst Pakistani women, acting both directly on cord-blood leptin levels and explaining 0.047 SD (81%) of the difference and also acting via cord-blood insulin levels, with that pathway explaining 0.019 SD (19%) of the difference.

### Discussion

We found positive linear associations of gestational fasting and postload glucose levels with cord-blood levels of insulin and leptin in both Pakistani and white British women. Pakistani infants had higher cord-blood leptin levels, indicating greater fat mass at birth, compared with white British infants. The
magnitude of this difference was modest, with Pakistani infants
having 16% higher levels of cord-blood leptin, equating to a
0.06SD difference. Given that the birthweight of Pakistani in-
fants was >250 g lower than that of white British infants, this
difference highlights a very different body composition between
the two ethnic groups. We found that this greater birth adiposity
was largely explained by higher gestational fasting glucose
levels in women of Pakistani origin compared with white British
women, which appears to act both directly on cord-blood leptin
levels and also via increased fetal insulin secretion (as measured
here by cord-blood insulin levels).

Glucose freely crosses the placenta; therefore, the amount
delivered to the developing fetus depends upon maternal
circulating levels [15–17]. Glucose levels are higher in Paki-
stanian than white British women, thus the developing fetuses of
Pakistani women will be relatively overnourished with glu-
cose, which will stimulate increased fetal pancreatic secretion
of insulin, which is a growth and fat deposition hormone. Our
regression and path analysis results support these pathways
and our original hypothesis that higher cord-blood leptin
levels (birth adiposity) in Pakistani infants is driven by higher
circulating gestational glucose levels in Pakistani women.

Table 2 Multivariable analyses of differences in cord-blood levels of leptin and insulin between Pakistani and white British mother–offspring pairs in all
participants and in participants without GDM

| Outcome                      | Modela | Mean difference (%) in outcome comparing Pakistani to white British (95% CI) |
|------------------------------|--------|-----------------------------------------------------------------------------|
|                              | Model 1: unadjusted | All participants n=1,415 | Participants without GDM n=1,285 |
| Cord-blood insulin level (%) | Model 1: unadjusted | 14 (6, 22) | 10 (1, 18) |
| Model 2: confounder          | 9 (0, 19) | 9 (0.4, 18) |
| Model 3: confounder and gestational fasting glucose level | 0 (−9, 11) | 5 (−4, 14) |
| Model 4: confounder and gestational postload glucose level | 5 (−5, 15) | 8 (−1, 17) |
| Model 5: confounder and gestational fasting and postload glucose levels | 0 (−10, 10) | 4 (−5, 13) |
| Cord-blood leptin level (%)  | Model 1: unadjusted | 13 (4, 21) | 9 (0.2, 19) |
| Model 2: confounder          | 16 (6, 26) | 12 (4, 21) |
| Model 3: confounder and gestational fasting glucose level | 7 (−3, 16) | 6 (−3, 14) |
| Model 4: confounder and gestational postload glucose level | 12 (3, 22) | 10 (1, 19) |
| Model 5: confounder and gestational fasting and postload glucose levels | 6 (−4, 16) | 5 (−4, 14) |
| Model 6: confounder and gestational fasting glucose and cord-blood insulin levels | 5 (−4, 14) | 4 (−4, 12) |

All results are mean differences of outcome comparing Pakistani mother–offspring pairs to White British mother–offspring pairs (reference group); positive values indicate higher levels of outcomes in Pakistani pairs and the null value for all results is 0

a Model 1, no adjustment; model 2, adjustment for potential confounding by maternal age, BMI, parity, smoking, education, gestational age and offspring
sex; model 3, as model 2 plus adjustment for gestational fasting glucose level; model 4, as model 2 plus adjustment for gestational postload glucose level;
model 5, as model 2 plus adjustment for gestational fasting and postload glucose levels; model 6, as model 2 plus adjustment for gestational fasting
glucose and cord-blood insulin levels

Fig. 3 Path analysis for differences in cord-blood leptin levels between
Pakistani and white British individuals (n=1415). The results are
standardised regression coefficients, with their 95% CIs, and are adjusted
for maternal age, parity and education, infant sex and gestational age.

They are interpreted as the adjusted change in outcome (box at the end of
the arrow head) in SD units per category (ethnicity) or per SD of the
exposure (arrow start). \( p<0.001 \) for all results except those indicated
\( p\geq0.50 \)
We undertook analyses both including and excluding women who met clinical criteria for GDM. In the Bradford Royal Infirmary, once identified, women with GDM will have been treated as clinically indicated with diet and exercise, metformin, or insulin to achieve and maintain euglycaemia. Treatment protocols are applied identically to women of all ethnicities. Women identified as having GDM are likely to have had lower glucose levels on average during the last trimester than if they had not been identified and treated; however, these women represent the upper end of the risk distribution and will have had higher glucose levels in earlier pregnancy and we felt it was important to examine relationships in the cohort with these women included as well as with them excluded. Overall, the mediation analyses were similar in cohorts that included or excluded women with GDM, with some minor differences between each stage of the regression and path analyses. GDM was more common in Pakistani than white British women, consistent with their higher levels of fasting and postload glucose, but in both ethnic groups most women were diagnosed with GDM on the basis of having elevated postload glucose levels. However, fasting glucose level was more strongly related to cord-blood levels of insulin and leptin than was postload glucose level, thus irrespective of inclusion or exclusion of those with GDM, the mediation by fasting glucose levels was similar.

To the best of our knowledge, this study is the first to examine the possibility that birth differences in cord-blood leptin levels between South Asian and white European populations is mediated by differences in gestational glycaemia. We used cord-blood insulin levels as a proxy for fetal insulin secretion and cord-blood leptin levels as a proxy for birth fatness. Maternal insulin does not cross the placenta and the fetus secretes insulin in relation to maternal levels of circulating glucose; the anabolic effects of insulin are then key for fetal fat deposition [15–17]. However, cord-blood insulin levels may also reflect insulin sensitivity at birth. Whilst it is difficult to determine how much of the variation in cord-blood insulin levels was due to offspring insulin sensitivity at birth, this sensitivity is likely to reflect only a small proportion of the variation, with variation largely reflecting fetal secretions [35, 36]. Leptin is found in fetal adipose tissue, and cord-blood leptin level is strongly positively related to fetal fat mass, explaining approximately 20–25% of variation in fat mass [26, 27], similar to the variation of birthweight that we found here. Whilst it is known that the placenta secretes leptin, the vast majority of this leptin is secreted into the maternal circulation, with less than 1% of cord-blood leptin estimated to be from the placenta/maternal circulation [37, 38]. Recent findings show that maternal gestational serum leptin level is not associated with cord-blood leptin level and is higher in women with a fetus with intrauterine growth restriction, whereas cord-blood leptin level is positively associated with birthweight [38]. These findings support cord-blood leptin level as reflecting fat mass at birth. Whilst we are unaware of any study examining whether cord-blood leptin level relates to fat mass with the same magnitude in European and South Asian infants, we found that the association of cord-blood leptin level with birthweight was the same in both groups, which is consistent with our previous more detailed study of these relationships [39].

Our study is observational and cannot prove causality. To assume the observed associations represent causal paths means assuming that measurement error between ethnicity and the mediators, and the mediators and cord-blood leptin, are minimal and/or are not correlated with each other [40]. Ethnicity was self-reported but verified by family tree information and country of birth of family members. Gestational glucose and cord-blood measurements were completed using valid procedures and with the laboratory staff unaware of participant characteristics. Claiming causal mediation would also assume that there is no residual confounding. Mendelian randomisation (using genetic variants that are known to be robustly associated with fasting glucose levels as proxies for its causal effect) has shown a causal effect of fasting glucose levels on birthweight, which together with the known physiology of pregnancy in relation to maternal glucose and fetal insulin, supports a causal effect of fasting glucose levels on fetal fatness. However, further replication of these findings and testing causal pathways through randomised controlled trials and/or Mendelian randomisation is warranted.

The South Asian participants in our study are all of Pakistani origin and we cannot assume that our results necessarily generalise to other South Asian populations. Likewise, the white British population may not be representative of all white European populations. However, other studies suggest similar differences in infant fat mass between Indian and white European infants [7–9].

Leptin regulates energy homeostasis by relaying information about the body’s energy and nutrient stores from the periphery to the brain [41]. However, adult humans in Western populations are largely resistant to this regulatory mechanism, with recent evidence suggesting that leptin resistance is present in children as young as 7 and 12 years of age [42, 43]. By contrast, two studies of 200 and 600 participants have shown that cord-blood leptin levels are inversely related to BMI at age 2–3 years [44, 45], with these findings interpreted as showing that there might be a period of leptin sensitivity in early human life. Both of these studies were conducted in European origin children and whether similar results would be seen in South Asian infants is yet to be determined. Furthermore, the substantial evidence of greater fatness, insulin resistance, type 2 diabetes and markedly higher leptin levels in South Asian children and adults compared with Europeans [1, 11–14, 20–25] argues against any leptin sensitivity in infancy having a lasting beneficial impact, but rather that higher leptin and fat mass levels are maintained in South Asians. Our results provide further evidence of the higher prevalence of hyperglycaemia in South Asian than white British
women and the potential adverse effects in terms of greater adiposity in the offspring of South Asian women. These findings suggest that current guidelines for the detection and control of hyperglycaemia and GDM in South Asian women should be rigorously applied. Selecting women for GDM testing on the basis of risk factors, as currently practised in most of the UK and many other countries, could miss nearly half of all cases of GDM in South Asian women because of failure to rigorously apply risk factor screening [46, 47]. Furthermore, to our knowledge, there is no randomised controlled trial evidence of the effect of treating GDM (or those at risk of GDM) specifically in South Asian women, despite their greater risk and differing lifestyles compared with white European women. In the most recent Cochrane systematic review, which included eight trials, none seem to have included any or sufficient South Asian women to determine effects in that population [48]. We were able to identify just one trial in South Asian women, which was a pilot randomised trial in India that aimed to compare the relative effects on GDM of two different types of insulin therapy [49].

In conclusion, we have shown that compared with white British infants, Pakistani infants have higher absolute cord-blood leptin levels, reflecting greater birth fat mass despite their lower birthweight. Importantly, we found that higher maternal gestational glucose levels explain this greater birth fat mass. Given the rapidly rising levels of diabetes in South Asian populations, these results warrant further exploration and, if replicated, the possibility of more aggressive identification and treatment of hyperglycaemia in pregnancy in South Asian women should be explored.

Acknowledgements We are grateful to all the families who took part in this study, the midwives for their help in recruiting them, the paediatricians and health visitors, and the Born in Bradford team that included interviewers, data managers, laboratory staff, clerical workers, research scientists, volunteers and managers.

Funding This study was supported by a British Heart Foundation Project Grant (PG09/036/27380).

DAL works in a unit that receives funding from the UK Medical Research Council (MRC) (MC_UU_12013/5). JWe was funded by a UK MRC Special Training Fellowship in Health of the Public and Health Services Research (MRCG0601712). No funding bodies had any role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript. The views expressed in this manuscript are those of the authors and not necessarily any funding body.

Duality of interest The authors declare that there is no duality of interest associated with this manuscript.

Contribution statement DAL contributed to designing and conducting the Born in Bradford (BiB) study, obtained funds for the study reported here, conceived, designed and developed the study presented here, wrote the statistical analysis plan and undertook analyses and wrote the first draft of the paper. She acts as guarantor. JWe contributed to designing and conducting the BiB study, obtained funds for the study reported here and contributed to data collection for this study. LF contributed to conducting the BiB study and managed and contributed to collection of the data used in this paper. SMN developed the study aim, conception and design. RSB contributed to designing and conducting the BiB study, obtained funds for the study reported here and contributed to developing the design of this study. DT contributed to designing and conducting the BiB study, obtained funds for the study reported here and developed the study aim and design.

DJF collected cord-blood insulin data and obtained funds for the study. JWr contributed to designing and conducting the BiB study and is the scientific director of BiB, he contributed to developing the concept and aim of this study and obtained funds for the study reported here. DCW obtained funds for the study reported here, developed the study aim and collected maternal gestational glucose data.

NS obtained funds for the study reported here and conceived and developed the study aim and design. JWe, LF, SMN, RSB, DT, DJF, JWr, DCW and NS made critical comments on drafts of the paper. All authors approved the final version of the paper.

Open Access This article is distributed under the terms of the Creative Commons Attribution License which permits any use, distribution, and reproduction in any medium, provided the original author(s) and the source are credited.

References

1. McKeigue PM, Pierpoint T, Ferrie JE, Marmot MG (1992) Relationship of glucose intolerance and hyperinsulinaemia to body fat pattern in south Asians and Europeans. Diabetologia 35:785–791
2. Yajnik CS, Yudkin JS (2004) The Y-Y paradox. Lancet 363:163
3. Ramachandran A, Snehala B, Vijay S (2004) Low risk threshold for acquired diabetogenic factors in Asian Indians. Diabetes Res Clin Pract 65:189–195
4. Ehtisham S, Crabtree N, Clark P, Shaw N, Barrett T (2005) Ethnic differences in insulin resistance and body composition in United Kingdom adolescents. J Clin Endocrinol Metab 90:3963–3969
5. Whincup PH, Gilg JA, Owen CG, Odoki K, Alberi KG, Cook DG (2005) British South Asians aged 13–16 years have higher fasting glucose and insulin levels than Europeans. Diabet Med 22:1275–1277
6. Nightingale CM, Rudnicka AR, Owen CG, Cook DG, Whincup PH (2011) Patterns of body size and adiposity among UK children of South Asian, black African–Caribbean and white European origin: Child Heart And Health Study in England (CHASE Study). Int J Epidemiol 40:33–44
7. Yajnik CS, Lubree HG, Rege SS et al (2002) Adiposity and hyperinsulinaemia in Indians are present at birth. J Clin Endocrinol Metab 87:5575–5580
8. Yajnik CS, Fall CH, Coya J et al (2003) Neonatal anthropometry: the thin–fat Indian baby. The Pune Maternal Nutrition Study. Int J Obes Relat Metab Disord 27:173–180
9. Krishnaveni GV, Hill JC, Veena SR et al (2000) Truncal adiposity is present at birth and in early childhood in South Indian children. Indian Pediatr 42:527–538
10. West J, Lawlor DA, Fairley L et al (2013) UK-born Pakistani-origin infants are relatively more adipose than white British infants: findings from 8704 mother–offspring pairs in the Born-in-Bradford prospective birth cohort. J Epidemiol Community Health 67:544–551
11. Simmons D (1992) Parity, ethnic group and the prevalence of type 2 diabetes: the Coventry Diabetes Study. Diabet Med 9:706–709
12. Dornhorst A, Paterson CM, Nicholls JS et al (1992) High prevalence of gestational diabetes in women from ethnic minority groups. Diabet Med 9:820–825
13. Makgoba M, Savvidou MD, Steer PJ (2012) An analysis of the interrelationship between maternal age, body mass index and racial origin in the development of gestational diabetes mellitus. BJOG 119: 276–282
14. Wong VW (2012) Gestational diabetes mellitus in five ethnic groups: a comparison of their clinical characteristics. Diabet Med 29:366–371
15. Lawlor DA (2013) The Society for Social Medicine John Pemberton Lecture 2011. Developmental overnutrition—an old hypothesis with new importance? Int J Epidemiol 42:7–29
16. Lawlor DA, Relton C, Sattar N, Nelson SM (2012) Maternal adiposity—a determinant of perinatal and offspring outcomes? Nat Rev Endocrinol 8:679–688
17. Catalano PM, Hauguel-De Mouzon S (2011) Is it time to revisit the Pedersen hypothesis in the face of the obesity epidemic? Am J Obstet Gynecol 204:479–487
18. The HAPO Study Cooperative Research Group (2008) Hyperglycemia and adverse pregnancy outcomes. N Engl J Med 358:1991–2002
19. The HAPO Study Cooperative Research Group (2009) Hyperglycemia and Adverse Pregnancy Outcome (HAPO) Study: associations with neonatal anthropometrics. Diabetes 58:453–459
20. Mather HM, Keen H (1985) The Southall Diabetes survey: prevalence of known diabetes in Asians and Europeans. BMJ 291:1081–1084
21. Bhopal R, Unwin N, White M et al (1999) Heterogeneity of coronary heart disease risk factors in Indian, Pakistani, Bangladeshi, and European origin populations: cross sectional study. BMJ 319:215–220
22. Barnett AH, Dixon AN, Bellary S et al (2006) Type 2 diabetes and cardiovascular risk in the UK south Asian community. Diabetesologia 49:2234–2236
23. Gholap N, Davies M, Patel K, Sattar N, Khunti K (2011) Type 2 diabetes and cardiovascular disease in South Asians. Prim Care Diabetes 5:45–56
24. Mather HM, Chaturvedi N, Fuller JH (1998) Mortality and morbidity from diabetes in South Asians and Europeans: 11-year follow-up of the Southall Diabetes Survey, London, UK. Diabet Med 15:53–59
25. Chan JC, Malik V, Jia W et al (2009) Diabetes in Asia: epidemiology, risk factors, and pathophysiology. JAMA 301:2129–2140
26. Clapp JF III, Kiess W (1998) Cord blood leptin reflects fetal fat mass. J Soc Gynecol Invest 5:300–303
27. Ashworth CJ, Hoggard N, Thomas L, Mercer JG, Wallace JM, Lea RG (2000) Placental leptin. Rev Reprod 5:18–24
28. Wright J, Small N, Raynor P et al (2013) Cohort Profile: The Born in Bradford multiethnic family cohort study. Int J Epidemiol 42:7–29
29. Office of National Statistics (2003) Ethnic group statistics: a guide for the collection and classification of ethnicity data. Office of National Statistics, London
30. WHO (1999) Definition, diagnosis and classification of diabetes mellitus and its complications. Report of a WHO Consultation. Part 1: Diagnosis and classification of diabetes mellitus. WHO Department of Noncommunicable Disease Surveillance, Geneva
31. International Association of Diabetes and Pregnancy Study Groups Consensus Panel, Metzger BE (2010) International Association of Diabetes and Pregnancy Study Groups recommendations on the diagnosis and classification of hyperglycemia in pregnancy. Diabetes Care 33:676–682
32. McDonald TJ, Perry MH, Peake RW et al (2012) EDTA improves stability of whole blood C-peptide and insulin to over 24 hours at room temperature. PLoS ONE 7:e42084
33. McConway MG, Johnson D, Kelly A, Griffin D, Smith J, Wallace AM (2000) Differences in circulating concentrations of total, free and bound leptin relate to gender and body composition in adult humans. Ann Clin Biochem 37:717–723
34. Royston P, Sauerbrei W (2003) Stability of multivariable fractional polynomial models with selection of variables and transformations: a bootstrap investigation. Stat Med 28:639–659
35. Catalano PM, Presley L, Minium J, Hauguel-de Mouzon S (2009) Fetuses of obese mothers develop insulin resistance in utero. Diabetes Care 32:1076–1080
36. Luo ZC, Delvin E, Fraser WD et al (2010) Maternal glucose tolerance in pregnancy affects fetal insulin sensitivity. Diabetes Care 33:2055–2061
37. Lepercq J, Challier JC, Guerre-Millo M, Cauzac M, Vidal H, Hauguel-de Mouzon S (2001) Prenatal leptin production: evidence that fetal adipose tissue produces leptin. J Clin Endocrinol Metab 86: 2409–2413
38. Pighetti M, Tommaselli GA, D’Elia A et al (2003) Maternal serum and umbilical cord blood leptin concentrations with fetal growth restriction. Obstet Gynecol 102:535–543
39. West JW, Fairley L, Sattar N, Whincup P, Lawlor DA (2014) Do ethnic differences in cord blood leptin levels differ by birthweight category? Findings from the Born in Bradford cohort study. Int J Epidemiol 43:249–254
40. Blakely T, McKenzie S, Carter K (2013) Misclassification of the mediator matters when estimating indirect effects. J Epidemiol Community Health 67:458–466
41. Zhang Y, Proenca R, Maffei M, Barone M, Leopold L, Friedman JM (1994) Positional cloning of the mouse obese gene and its human homologue. Nature 372:425–432
42. Fleisch AF, Agarwal N, Roberts MD et al (2007) Influence of serum leptin on weight and body fat growth in children at high risk for adult obesity. J Clin Endocrinol Metab 92:948–954
43. Salbe AD, Weyer C, Lindsay RS, Ravussin E, Tataranni PA (2002) Assessing risk factors for obesity between childhood and adolescence: I. Birth weight, childhood adiposity, parental obesity, insulin, and leptin. Pediatrics 110:299–306
44. Brunner S, Smidt D, Huttinger K et al (2014) Effect of reducing the n-6/n-3 fatty acid ratio on the maternal and fetal leptin axis in relation to infant body composition. Obesity (Silver Spring) 22:217–224
45. Mantzoros CS, Rifas-Shiman SL, Williams CJ, Fargnoli JL, Kelesidis T, Gillman MW (2009) Cord blood leptin and adiponectin as predictors of adiposity in children at 3 years of age: a prospective cohort study. Pediatrics 123:682–689
46. Wagasaarachchi PT, Fernando L, Premachandra P, Fernando DJS (2011) Screening based on risk factors for gestational diabetes in an Asian population. J Obstet Gynaecol 21:32–34
47. Farrar D, Fairley L, Lawlor DA, Tuffnell D, Whitelaw D, Wright J (2014) Evaluation of the impact of universal testing for gestational diabetes mellitus on maternal and neonatal health outcomes: a retrospective analysis. BMC Pregnancy Childbirth. doi:10.1186/1471-2393-14-137
48. Alwan N, Tuffnell DJ, West J (2009) Treatments for gestational diabetes. Cochrane Database Syst Rev (3): CD003395. doi:10.1002/14651858.CD003395.pub2
49. Balaji V, Balaji MS, Alexander C et al (2010) Premixed insulin aspart 30 (BiAsp 30) vs premixed human insulin 30 (BHI 30) in gestational diabetes mellitus—a pilot study. J Assoc Physicians India 58:99–101