**in a multi-ethnic UK birth cohort: associations with child cognitive and educational development**

**Running title:** Maternal iodine & child development

Diane E Threapleton1\*, Charles JP Snart1\*, Claire Keeble1,2, Amanda H Waterman3, Elizabeth Taylor1, Dan Mason4, Stephen Reid5, Rafaq Azad4, Liam JB Hill3, Sarah Meadows6,7, Amanda McKillion6,7, Nisreen A Alwan8,9, Janet E Cade10, Nigel AB Simpson11, Paul M Stewart12, Michael Zimmermann13, John Wright4, Dagmar Waiblinger4, Mark Mon-Williams3, Laura J Hardie1†, Darren C Greenwood1,2†

\*Joint first authors

†Joint senior authors

1. Leeds Institute of Cardiovascular & Metabolic Medicine, School of Medicine, University of Leeds, Leeds LS2 9JT

2. Leeds Institute for Data Analytics, University of Leeds

3. School of Psychology, University of Leeds, Leeds, LS2 9JT, UK

4. Bradford Institute for Health Research, Bradford Teaching Hospitals NHS Foundation Trust, Bradford, BD9 6RJ, UK

5. Earth Surface Science Institute, School of Earth and Environment, University of Leeds, Leeds, LS2 9JT, UK

6. Elsie Widdowson Laboratory, University of Cambridge, Cambridge, CB1 9NL, UK

7. NIHR Nutritional Biomarker Laboratory, MRC Epidemiology Unit, University of Cambridge, Clifford Allbutt Building, Hills Road, Cambridge, CB2 0AH, UK

8. School of Primary Care and Population Sciences, Faculty of Medicine, University of Southampton, Southampton General Hospital, Southampton, SO16 6YD, UK

9. NIHR Southampton Biomedical Research Centre, University of Southampton and University Hospital Southampton NHS Foundation Trust

10. Nutritional Epidemiology Group, School of Food Science & Nutrition, University of Leeds, Leeds, LS2 9JT, UK

11. Nigel AB Simpson Division of Women’s and Children’s Health, School of Medicine, University of Leeds, Leeds, LS2 9JT, UK

12. Faculty of Medicine and Health, University of Leeds, Leeds, LS2 9JT, UK

13. Laboratory for Human Nutrition, Institute of Food, Nutrition and Health, ETH Zurich, 8092 Zürich, Switzerland

**Corresponding authors:**

Darren C Greenwood and Laura J Hardie

Leeds Institute of Cardiovascular & Metabolic Medicine,

School of Medicine,

University of Leeds,

Leeds LS2 9JT

United Kingdom

d.c.greenwood@leeds.ac.uk l.j.hardie@leeds.ac.uk

**Synopsis**

Study question: What is the association between maternal iodine status, child development and educational outcomes?

What’s already known: Maternal iodine requirements increase during pregnancy to supply thyroid hormones critical for fetal neurodevelopment. Severe iodine insufficiency may result in poorer cognitive outcomes but current evidence with mild-to-moderate insufficiency is sparse and inconsistent.

What this study adds: There was little evidence maternal iodine status was associated with poorer neurodevelopmental outcomes, or with important objective educational outcomes at ages 4 to 7.

**Abstract**

Background: Maternal iodine requirements increase during pregnancy to supply thyroid hormones critical for fetal neurodevelopment. Iodine insufficiency may result in poorer cognitive or child educational outcomes but current evidence is sparse and inconsistent.

Objectives: To quantify the association between maternal iodine status and child educational outcomes.

Methods: Urinary iodine concentrations (UIC) and iodine/creatinine ratios (I:Cr) were measured in 6971 mothers at 26-28 weeks’ gestation participating in the Born in Bradford cohort. Maternal iodine status was examined in relation to child school achievement (Early Years Foundation Stage (EYFS), Phonics, and Key Stage 1 (KS1)), other learning outcomes, social and behavioural difficulties, and sensorimotor control in 5745 children aged four to seven years.

Results: Median (inter-quartile range) UIC was 76 µg/L (46, 120) and I:Cr was 83 µg/g (59, 121). Overall, there was no strong or consistent evidence to support associations between UIC or I:Cr and neurodevelopmental outcomes. For instance, predicted EYFS and Phonics scores (primary outcomes) at the 25th vs. 75th I:Cr percentiles (99% confidence intervals) were similar, with no evidence of associations: EYFS scores were 32 (99% CI 31, 33) and 33 (99% CI 32, 34), and Phonics scores were 34 (99% CI 33, 35) and 35 (99% CI 34, 36), respectively.

Conclusions: In the largest single study of its kind, there was little evidence of detrimental neurodevelopmental outcomes in children born to pregnant women with iodine insufficiency as defined by World Health Organization-outlined thresholds. Alternative functional biomarkers for iodine status in pregnancy and focused assessment of other health outcomes may provide additional insight.

**Keywords**

Iodine, child development, cognition, pregnancy, nutrition, deficiency, pregnancy, Born in Bradford

**Social Media Quote**

The mothers in our large cohort of pregnant women in the UK had insufficient iodine status according to World Health Organization definitions but there was no evidence this made any difference to their children’s development or school qualifications.

(Supported by Figure 1a and/or Figure 1b)

**Word count**

Abstract: 254 words. Main text: 3337 words.

**Background**

Iodine is an essential mineral required for thyroid hormone production, supporting normal metabolic processes throughout the life course.1 During pregnancy, iodine demands increase to support normal fetal development and compensate for increased renal iodine clearance.2 Because thyroid hormones are necessary for normal neuronal migration and myelination during brain development, severe maternal iodine deficiency can potentially hinder child growth and development, including psychomotor and neurological development.1,3 Irreversible mental retardation and neurological abnormalities can result from hypothyroxinaemia during critical developmental periods.2

Pregnant women in countries without iodisation or supplementation programmes are at particular risk of deficiency if they have low intakes of dairy, fish or meat.3 Whilst improved general nutrition has resulted in a diminishing prevalence of iodine deficiency disorders in the UK and Western Europe,1 iodine intake remains potentially inadequate, particularly in vulnerable groups. The World Health Organization defines iodine insufficiency in pregnant populations as median urinary iodine concentration (UIC) <150 µg/L, with several studies reporting insufficient iodine status in pregnant women at this concentration, but there is little evidence for the functional importance of this threshold.1,5

A small number of observational studies in developed countries have highlighted potential negative consequences of maternal iodine insufficiency for subsequent child cognition.4-12 In each case, associations were observed with only some outcome measures that were not consistent across studies. Furthermore, most previous work used existing WHO thresholds or assumed linear associations to examine neurodevelopmental outcomes4-7,10,11 rather than exploring if whether any thresholds exist for cognitive outcomes, or where they are. Further inconsistency was identified by a recent randomized controlled trial of iodine supplementation in mildly iodine-deficient pregnant women, which found no evidence of impact on offspring development at 5 years.13

This study therefore reports on maternal iodine concentrations during pregnancy using a large and well-characterised multi-ethnic cohort of British women, exploring the potential for threshold effects. We apply a comprehensive range of key neurodevelopmental domains in their offspring including objective measures of school achievement, standardised assessments of sensorimotor control and literacy, and teacher-reported assessments of emotional and behavioural development.

**Methods**

Cohort selection

All babies born to women who agreed to participate in the Born in Bradford cohort study were eligible,14 recruiting 12453 women with 13963 pregnancies between 2007 and 2010. Spot urine samples were collected at 26-28 weeks' gestation from 6644 women (see eFigure 1). The current study protocol has been registered at clinicaltrials.gov NCT03552341. The large study size provides sufficient power for detecting associations even of a potentially modest size (see protocol).

Exposure

Participants provided urine samples after overnight fasting at routine antenatal clinics for oral glucose tolerance tests. Samples were centrifuged, aliquoted and barcode-tagged before freezing and storage at -80 oC at Bradford Royal Infirmary and subsequent transfer to Leeds Institute of Cardiovascular and Metabolic Medicine, University of Leeds. Urinary 127iodine concentration (µg/L) was estimated using inductively coupled plasma mass spectrometry (ICP-MS) (Thermo iCAP Q, Hemel Hempstead, UK). This methodology is accredited by the Centers for Disease Control and Prevention (CDC) Ensuring the Quality of Urinary Iodine Procedures (EQUIP) standardisation programme.15 To improve accuracy in measuring iodine status and account for urine dilution, we used the iodine-to-creatinine ratio (I:Cr), which has been shown to better reflect 24-hour urine excretion than UIC in pregnant women.16 A standard microplate assay utilising the Jaffe reaction was employed for evaluating creatinine concentrations. For inter-laboratory comparison, a subsample of the BiB urine samples were provided to the MRC Elsie Widdowson Laboratory, University of Cambridge, who conduct iodine analysis for the UK National Diet and Nutrition Survey (NDNS). Further details of laboratory methods are reported in Supplementary Methods.

Outcomes

The primary outcomes were Early Years Foundation Stage (EYFS) profile (age 4-5), Phonics score (age 5-6), and Key Stage 1 (KS1) school assessments (age 6-7). These assessments are reported to the local authority and data routinely reported to BiB. Additional learning, behavioural and sensorimotor assessments were also conducted during EYFS (age 4–5).17

The EYFS profile is a UK-wide, teacher-led, statutory evaluation of development in five prime areas of learning (personal, social and emotional development, physical development, and communication and language) and two specific areas of learning (mathematics and literacy).18 These are derived from assessments of achievement in 17 separate Early Learning Goals (ELGs).18 Children were scored as emerging (1), at expected standard (2) or exceeding standard (3) for each ELG, with total scores ranging from 17–51. Children were dichotomised as being “at or above the expected level” or “emerging”, across all and within each domain.18 The national EYFS assessment was updated in 2013, with scores being incomparable across pre- and post-2013 versions, leading to the exclusion of results from 303 children using the older version.

The Phonics assessment focuses on reading of 20 real and 20 pseudo words. Where children repeated the test the next year, results from the first attempt were used. KS1 attainment is measured within key domains (English, Reading, Writing, Mathematics and Science). The national KS1 assessment methods changed in 2016 during the study and results from the different versions were therefore analysed separately, with the KS1 ‘English’ sub-assessment discontinued after 2016.

Further developmental outcomes were collected for a subgroup of children.21 Receptive vocabulary was assessed using the British Picture Vocabulary Scale (BPVS) (2nd Edition). Letter Identification used a sub-test of the Woodcock Reading Mastery Test battery. The Strengths and Difficulties Questionnaire (SDQ) assessed child mental health difficulties as a score out of 40, combining emotional and peer problems (internalising) with conduct and hyperactivity problems (externalising)22 (Web Appendix, p2).

Children’s sensorimotor control was measured using the Clinical Kinematic Assessment Tool (CKAT), a tablet-based computerised assessment of children’s uni-manual tracking, aiming and steering performance (Supplementary Methods).

Statistical analysis

To account for variation in urine dilution23 the standardised measure of I:Cr was the primary exposure used, though UIC was also examined. Potential associations between iodine status and neurodevelopment were examined using multiple linear or logistic regression. All models used robust (Sandwich) estimates of variance, taking account of sibling clusters.24

We examined evidence for non-linear associations by fitting restricted cubic splines with four knots placed at 5th, 35th, 65th and 95th percentiles.25 Adjusted predicted outcomes were plotted for a standard lower-risk cohort participant (primiparous, white ethnicity, English as the dominant language, employed and not materially deprived, non-smoker, non-alcohol drinker, no pregnancy complications, with mean BMI, age and gestation). This was achieved by centring continuous covariates about the cohort mean and specifying the reference group for categorical variables. The Wald test was used to assess the overall contribution of iodine status to each model.

Adjustment for confounding was informed by a directed acyclic graph (eFigure 2). All models were adjusted for potential confounders and competing exposures (Supplementary Methods): maternal age, socioeconomic and educational status (eTable 2), ethnicity, season, smoking in pregnancy, drinking alcohol in pregnancy, pregnancy complications (gestational diabetes, hypertension, pre-eclampsia), early pregnancy body mass index (BMI), parity, child's sex, length of gestation and speaking English as an additional language.

To mitigate against multiple testing, 99% CIs were used throughout. Stata version 15·1 (StataCorp) was used for data preparation and analysis.

Missing data

Information was incomplete for some covariates, ranging from <1% missing for ethnicity, between 3-7% for clinical complications such as preeclampsia, pregnancy-induced hypertension, pre pregnancy hypertension and gestational diabetes, and between 10-12% for smoking status, alcohol use, English as an additional language, socioeconomic status & education and BMI (eTable 3). We imputed for missing covariates using multiple imputation by chained equations, with predictive mean matching for continuous variables and logistic regression for categorical variables. Separate imputation models were used for each outcome and exposure, including the outcome and all covariates in each imputation model, with 100 imputed datasets (Supplementary Methods).

Sensitivity analysis

Sensitivity analysis was conducted to confirm no substantial differences with results from unimputed data. Sensitivity analyses were also conducted additionally adjusting for total fish intake, consuming five portions of fruit and vegetables per day, and having ever breastfed, available in a subsample of the cohort (n=2776).

Further sensitivity analyses were conducted to assess robustness of results: (i) excluding extreme iodine concentrations >3 standard deviations (SD) from the mean on the log scale) (n=42), (ii) excluding women with pregnancy complications (gestational diabetes, hypertension or preeclampsia (n=913), and (iii) excluding users of iodine-containing supplements (n=1040). Evidence of effect-modification by ethnicity (White or Pakistani ethnic background), maternal socioeconomic and education category (more deprived and less educated vs less deprived and more educated), or child’s sex, were formally tested by including an interaction term in the model (Supplementary Methods).

Ethics approval

Ethical approval for BiB was granted by Bradford Research Ethics Committee (07/H1302/112).

**Results**

Study participants

Of 12453 women with 13776 pregnancies recruited into the cohort, 6644 women (53%) provided 6977 urine samples at 26-28 weeks' gestation. In total, 6971 samples were successfully analysed for iodine and creatinine from 6639 women with 7013 children. Women who provided urine samples were broadly similar to those who did not (eTable 4). The mean age of participants was 27 years (SD 6) and mean BMI was 26 kg/m2 (SD 6) (Table 1). Maternal characteristics and child outcomes are presented by maternal urinary I:Cr in Table 1 and 2. At least one outcome measure was collected or reported for 5745 (82%) children. Because of incomplete covariate data, subsequent results are based on the imputed datasets.

Iodine status

The median (inter-quartile range (IQR)) UIC for all samples was 76 µg/L (45 to 120) with 29% below 50 µg/L (Table 1), characterising this population of pregnant women as iodine insufficient by WHO criteria.26 In White European women, the median UIC (82 µg/L, IQR 49 to 127) was higher than women of Pakistani (73 µg/L, IQR 42 to 115), or other ethnic background (72 µg/L, IQR 42 to 111) (eTable 5). Median (IQR) I:Cr was 83 µg/g (59 to 121) and women with lower I:Cr were more likely to have lower socioeconomic and educational backgrounds (Table 1).

National school assessments

There was some evidence of higher maternal I:Cr associated with higher total EYFS score for I:Cr less than approximately 150 µg/g, though wide confidence intervals allowed the possibility of no association (Figure 1a). For a typical mother with lower-risk characteristics (defined above) the EYFS scores (99% CIs) at the 25th (59 µg/g), 50th (83 µg/g), and 75th (121 µg/g) I:Cr percentiles were predicted to be 32 (99% CI 31, 33), 33 (99% CI 32, 34), and 33 (99% CI 32, 34) µg/g, respectively (eTable 6). Sensitivity analyses using complete data only and excluding extreme I:Cr values (n = 42) yielded similar estimates (eTable 6, eFigure 3a-b). There was evidence for the association between I:Cr and EYFS score differed between boys and girls (pinteraction=0·005) with a positive relationship in girls but an inverse relationship in boys (eFigure 3c-d).

There was little evidence of an association between I:Cr and Phonics scores (Figure 1b) or in subgroups (child's sex, ethnic group or deprivation and educational group) (eFigure 4), though the general trends were consistent with that seen with EYFS scores. Estimates were similar after excluding I:Cr outliers (n=42) or mothers with complications in pregnancy (n=913)(eFigure 4a-b). There was no evidence of any association between I:Cr and achieving the EYFS or Phonics standard (Figure 1c-d) or achieving the EYFS standard in any sub-domain (eFigure 5).

Pre-2016 KS1 reading was the only KS1 test showing evidence of an association with maternal I:Cr (Figure 2a and 2b). For a typical mother with lower-risk characteristics (previously defined) the % achieving the pre-2016 KS1 reading standard (99% CIs) at the 25th, 50th and 75th I:Cr percentiles were estimated as 85 (99% CI 70, 93), 87 (99% CI 73, 94), and 84 (99% CI 68, 93) µg/g respectively (eTable 6). There was no evidence that this association differed by child's sex, ethnic background or maternal deprivation and education (eFigure 6). For other KS1 assessments, the pre-2016 assessments for maths, science and writing reflected the general trends seen with pre-2016 KS1 reading (Figures 2d, 2f and 2h), but wide confidence intervals included the possibility of no association. There was no evidence of any association with any of the post-2016 assessments (Figures 2c, 2e, 2g and 2i), sensitivity analyses, or of differences between subgroups (data not shown).

Assessment for learning, sensorimotor control and behavioural difficulties

SDQ scores (total, internalising and externalising) were not associated with I:Cr in any analyses (eFigure 7a-c), nor when SDQ scores were dichotomised (SDQ score ≥12, indicating slightly raised or high)(eFigure 7d). There was some evidence that higher I:Cr predicted fewer difficulties in White Europeans than in children with Pakistani ethnic backgrounds (pinteraction=0.005) (eFigure 8c-d), but no evidence of a difference between males and females (pinteraction=0.847) (eFigure 8a-b).

There was no evidence of any association between I:Cr and Letter identification, receptive vocabulary, or sensorimotor control (eFigure 9). In sensitivity analyses, additionally adjusting for diet (fruit, vegetable and fish intake) or breastfeeding gave similar estimates (data not shown). Iodine measured as UIC rather than I:Cr was not found to be associated with any outcomes (eTable 6).

**Comment**

Principal findings

In this UK cohort, where dietary fortification or supplementation of iodine was not routine, there was little evidence of associations between iodine status measured at 26-28 weeks' gestation and most child educational, learning or sensorimotor assessments at ages 4–7 years. The single exception to this was in just one of nine KS1 outcomes (pre-2016 KS1 reading). However, whilst uncertain, trends in EYFS scores, Phonics and pre-2016 KS1 assessments were all consistent and there was some evidence of modest positive associations between I:Cr and the EYFS, Phonics and SDQ scores in some sensitivity analyses and subgroups.

Strengths of the study

Key strengths in our methodology include objective national measures of educational achievement (outcomes) and robust sample analysis using CDC-endorsed and validated urinary iodine assessment methods (exposure).15 We also corrected UIC for creatinine concentrations to account for important differences in urine dilution.19 Our modelling of potential nonlinear associations permits identification of any important I:Cr thresholds, though we did not observe any consistent evidence of a threshold effect in this setting. Further strengths include a population with comparatively low iodine status for a developed country allowing a wide range of iodine exposure, and from a multi-ethnic community allowing a wide range of dietary intakes. Our findings support the suggestion that people from South Asian backgrounds are more likely to have insufficient iodine.

Limitations of the data

There are limitations in assessing iodine status from single spot urine samples, including high day-to-day variation in iodine intake not reflecting a usual iodine status,27 any rapid changes during pregnancy or brief periods of insufficiency. However, this method is widely used in population studies and is considered sufficient for characterising iodine status in populations.26 Furthermore, urinary iodine excretion has been found to reflect recent iodine intake27 and all samples in this study were collected after overnight fasting and at a similar time of day. Urine samples were unavailable for a proportion of mothers and there was incomplete follow-up for some developmental outcomes. However, this was mostly because of funding constraints rather than non-response, and women providing samples were similar to those who did not, so unlikely to introduce bias. Our multi-ethnic community may not be representative of the UK, but this affords a range of exposures providing greater opportunity to identify associations with lower maternal iodine status as well as high.

Interpretation

Other cohorts previously reported evidence for associations between low iodine status and poorer neurodevelopmental measures, but findings are not consistent across different studies using similar outcomes, or across different outcomes within the same study.5,7-9 Whilst there remains insufficient evidence of any strong associations within our cohort, our findings are more consistent with studies suggesting weak associations in more literacy-based outcomes, language, verbal IQ and reading, rather than numeracy or motor skills.4,5,7-9 Some of the inconsistencies reported across studies may in part reflect different iodine measures. 24h urinary iodine excretion is the gold standard measure for iodine status but may not be feasible in large cohort studies, with spot sample UIC and I/Cr measures preferred. I/Cr minimizes the variations caused by urine volume differences and dilution in pregnancy, and better reflects 24-hour iodine excretion and circulating iodine levels during pregnancy and postpartum.23,28 UIC may also increase the apparent prevalence of iodine deficiency compared with I/Cr measurement.18

Of existing comparable studies reporting urinary iodine status in pregnancy and neurodevelopmental outcomes in children, this is the largest single cohort by some margin.4-9,11,12 Several other UK-based studies in pregnant women have also suggested iodine insufficiency according to WHO criteria,26 but the BiB cohort had lower iodine status than most (eTable 1). This provided a wider range of lower levels of exposure over which to detect any trends and allows the BiB cohort to contribute further to the evidence base.

Despite plausible mechanistic evidence for associations between maternal iodine status, thyroid hormones and fetal neurodevelopment,29 the evidence from our cohort of mothers does not provide strong evidence to support the hypothesis that insufficient iodine in pregnancy results in substantive adverse educational and cognitive outcomes in the child, within the observed range of mild-to-moderate iodine insufficiency.

It is not possible to identify whether any differences observed between boys and girls (EYFS) are because of biological differences provoking gendered behaviour patterns, or whether different expectations are responsible.30 Previous research using measures such as IQ tests aim to predict future educational outcomes. Whilst it can be argued that IQ tests may be more sensitive, we argue that it is ultimately educational outcomes that matter more to the future potential of the individual child and for the nation. We therefore consider it a strength to have linked and reported actual National measures of educational achievement rather than proxy measures. The Phonics and KS1 tests are substantively objective assessments, less vulnerable to response bias, which may otherwise account for the observed differences.

The association between higher maternal I:Cr and higher probability of achieving pre-2016 KS1 reading standards was not seen in with the post-2016 assessment, despite a larger sample of children, possibly reflecting different assessment criteria. Evidence for associations between I:Cr and raised SDQ scores were inconsistent across analyses and ethnic groups. It is possible that the SDQ score is more sensitive at higher values, or the subjective nature of the assessment may allow some unconscious bias, leading to differences between ethnic groups. Additionally, despite carefully controlling for important potential confounders, it we cannot rule out the potential for residual confounding.

The absence of consistent and strong associations may reflect a relatively small contribution of maternal iodine to overall child neurodevelopment compared with other influences, such as child diet, environmental or social factors. Alternatively, there may be misclassification of outcomes or measurement error in assessments or other measurement error in estimating iodine concentration from a single spot urine sample, though all samples were handled consistently and ICP-MS used throughout. Iodine at 26-28 weeks' gestation may be outside of a critically important time window,14 with a previous meta-analysis reporting associations between verbal IQ, but not non-verbal, with iodine status in first trimester.14 However, there is evidence that maternal iodine status does not change markedly through pregnancy,31,32 so women with lower iodine status in mid-pregnancy were also likely to have lower iodine status earlier in pregnancy, and any impact of changes would be smoothed somewhat by any storage in the thyroid. It may also be that urinary iodine is not the most sensitive marker of any risk from iodine inadequacy.1

Conclusions

In a population with relatively low iodine intake, maternal iodine status at 26-28 weeks' gestation was not consistently associated with most educational, learning, behavioural or sensorimotor outcomes in the majority of children aged four to seven years. Whilst some previous studies have found evidence for associations,4,5,7-9 findings have been inconsistent both within, and across, studies. Our study suggests caution when linking non-severe iodine insufficiency and neurodevelopmental outcomes. Despite this, there remains scope for examining maternal iodine levels before conception and changes throughout pregnancy in such contexts, and the relationship with objectively-measured and specific neurodevelopmental outcomes in children. Identification of additional biomarkers that better characterise and accurately measure longer-term iodine status in individuals, alongside exploration of timing of samples, may also resolve inconsistencies in the evidence.

**Acknowledgements**

Born in Bradford is only possible because of the enthusiasm and commitment of the Children and Parents in BiB. We are grateful to all the participants, health professionals and researchers who have contributed to Born in Bradford. The authors thank Dr. Gillian Santorelli for advice relating to BiB data. This publication is independent research funded by the National Institute for Health Research (NIHR) Policy Research Programme (Assessing iodine status and associated health outcomes in British women during pregnancy, PR-R10-0514-11004). The views expressed are those of the authors and not necessarily those of the NIHR or the Department of Health and Social Care.

**Funding**

This publication is independent research funded by the National Institute for Health Research (NIHR) Policy Research Programme (Assessing iodine status and associated health outcomes in British women during pregnancy, PR-R10-0514-11004). The Born in Bradford study presents independent research commissioned by the National Institute for Health Research Collaboration for Applied Health Research and Care (NIHR CLAHRC) and the Programme Grants for Applied Research funding scheme (RP-PG-0407-10044). The views expressed in this publication are those of the authors and not necessarily those of the NHS, the NIHR or the Department of Health and Social Care.

**Data Sharing and Data Accessibility**

Born in Bradford welcome collaboration with other researchers. Requests for existing data and biological samples will be reviewed, prioritised and authorised by the BiB Executive Group. Potential collaborators should complete an outline proforma available on the Born in Bradford website (borninbradford.nhs.uk) and submit to the BiB Director.

**Figure legends**

**Figure 1 Estimated scores and the estimated % of children achieving the EYFS and Phonics standards across the range of maternal urinary I:Cr concentrations**

Abbreviations: ELG early learning goals; EYFS Early Years Foundation Stage.

Histograms illustrate the distribution of iodine concentrations and although the figures are curtailed at 300 µg/g, the splines (solid line) and 99% CIs (dashed lines) were drawn using data from all participants. Splines were drawn after adjustment for maternal age, ethnic group, season, socioeconomic status, BMI, parity, smoking, alcohol, complications in pregnancy, gestation length, child's sex, and speaking English as an additional language.

The spline position in these figures illustrates the predicted estimate for typical participants (primiparous, white ethnic background, do not speak English as an additional language, 'Employed and not materially deprived', did not smoke, drink or experience complications in pregnancy and have average BMI, age and gestation length).

**Figure 2 Estimated % of children achieving the Key Stage 1 standards across the range of maternal I:Cr concentrations**

Adjustments and figure details are as in Figure 1. The KS1 standard before 2016 is defined as 'Working securely at level 2B or beyond’. From 2016 the standard is defined as ‘Working at the expected standard or beyond’.

**Table 1 Maternal characteristics according to urinary iodine to creatinine ratio, n= 6971**

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
|  | | |  | **Maternal urinary iodine to creatinine ratio (fifths)** | | | | |
|  | | | **All participants** | **1 - lower** | **2** | **3** | **4** | **5 - higher** |
| I:Cr (µg/g), median (range) | | | 83 (1 to 2283) | 45 (1 to 54) | 64 (54 to 73) | 83 (73 to 95) | 111 (92 to 136) | 174 (136 to 2283) |
| n | | | 6971 | 1395 | 1394 | 1394 | 1394 | 1394 |
| I:Cr (µg/g), geometric mean (99% CI) | | | 86·1 (84·6, 87·6) | 41·9 (41·1, 42·7) | 63·7 (63·4, 64·1) | 83·2 (82·8, 83·7) | 111·7 (111·0, 112·5) | 190·4 (186·2, 194·7) |
| I:Cr (µg/g), median (IQR) | | | 83·1 (59·3, 121·0) | 44·8 (37·6, 50·1) | 64·0 (59·3, 68·7) | 83·1 (77·8, 89·1) | 110·8 (102·5, 121·0) | 174·0 (151·5, 218·2) |
| UIC (µg/L), geometric mean (99% CI) | | | 70·8 (69·2, 72·5) | 41·1 (39·1, 43·2) | 55·6 (53·1, 58·2) | 72·5 (69·4, 75·7) | 84·4 (80·9, 88·1) | 127·6 (121·7, 133·7) |
| UIC (µg/L), median (IQR) | | | 76·2 (44·7, 120·2) | 46·6 (22·2, 71·2) | 61·8 (36·3, 91·3) | 80·9 (51·0, 114·2) | 92·9 (55·4, 133·6) | 137·8 (83·6, 207·9) |
| Age (years), mean (SD) | | | 27·2 (5·6) | 26·4 (5·5) | 26·7 (5·5) | 27·6 (5·6) | 27·4 (5·7) | 28·0 (5·5) |
| BMI (Kg/M2), mean (SD) | | | 25·9 (5·5) | 26·7 (5·9) | 26·3 (5·6) | 25·9 (5·5) | 25·4 (5·0) | 25·1 (5·1) |
| Educational achievement1, n (%) | | |  |  |  |  |  |  |
|  | <5 GSCE equivalent | | 1271 (18) | 325 (23) | 266 (19) | 255 (18) | 212 (15) | 213 (15) |
|  | 5 GSCE equivalent | | 1827 (26) | 416 (30) | 398 (29) | 342 (25) | 352 (25) | 319 (23) |
|  | A-level equivalent | | 932 (13) | 167 (12) | 176 (13) | 202 (14) | 199 (14) | 188 (13) |
|  | Higher than A-level | | 1707 (24) | 242 (17) | 295 (21) | 362 (26) | 377 (27) | 431 (31) |
|  | Don’t know/ other | | 1234 (18) | 245 (18) | 259 (19) | 233 (17) | 254 (18) | 243 (17) |
| Socio-economic status | | |  |  |  |  |  |  |
|  | Least deprived and most educated | | 1312 (21) | 179 (15) | 221 (18) | 294 (24) | 279 (23) | 339 (27) |
|  | Employed and not materially deprived | | 1291 (21) | 191 (16) | 250 (20) | 264 (21) | 306 (25) | 280 (22) |
|  | Employed and no access to money | | 961 (16) | 184 (15) | 213 (17) | 183 (15) | 189 (15) | 192 (15) |
|  | Receiving benefits and not materially deprived | | 1174 (27) | 430 (35) | 362 (29) | 304 (25) | 284 (23) | 294 (23) |
|  | Most economically deprived | | 954 (15) | 240 (20) | 191 (15) | 192 (16) | 181 (15) | 150 (12) |
| Ethnic background, n (%) | | |  |  |  |  |  |  |
|  | White European | | 3010 (43) | 472 (34) | 584 (42) | 623 (45) | 645 (47) | 686 (49) |
|  | Pakistani | | 2946 (43) | 678 (49) | 626 (45) | 573 (41) | 560 (40) | 509 (37) |
|  | Other (Black, Indian, mixed, other) | | 970 (14) | 231 (17) | 179 (13) | 186 (13) | 180 (13) | 194 (14) |
| Speaks English as an additional language | | | 2653 (43) | 627 (51) | 593 (47) | 508 (41) | 469 (37) | 456 (37) |
| Health and lifestyle in pregnancy | | |  |  |  |  |  |  |
|  | Complication in pregnancy3, n (%) | | 896 (13) | 168 (12) | 178 (13) | 174 (12) | 186 (13) | 190 (14) |
|  | Drank any alcohol, n (%) | | 1323 (19) | 217 (16) | 269 (19) | 291 (21) | 265 (19) | 281 (20) |
|  | Smoked, n (%) | | 1012 (15) | 188 (13) | 200 (14) | 225 (16) | 191 (14) | 208 (15) |
|  | Used any supplements, n (%) | | 1398 (20) | 120 (9) | 167 (12) | 255 (18) | 320 (23) | 536 (38) |
|  | Used iodine-containing supplement, n (%) | | 1040 (15) | 65 (5) | 106 (8) | 178 (13) | 235 (17) | 456 (33) |
|  | White fish intake2 (g/d), mean (SD) | | 21·0 (26·9) | 19·5 (28·1) | 18·3 (24·0) | 25·5 (29·5) | 21·9 (26·8) | 22·5 (25·7) |
|  | Oily fish intake2 (g/d), mean (SD) | | 1·4 (3·9) | 1·1 (3·7) | 1·0 (3·0) | 1·4 (3·9) | 1·9 (4·5) | 1·8 (4·1) |
|  | Total fish intake2 (g/d), mean (SD) | | 23·9 (29·5) | 22·2 (31·2) | 20·5 (26·2) | 25·4 (31·8) | 24·9 (29·2) | 26·4 (28·3) |
|  | Eats 5 portions of fruit and vegetables per day2, n (%) | |  |  |  |  |  |  |
|  |  | Always | 441 (19) | 77 (15) | 87 (19) | 96 (19) | 95 (21) | 86 (19) |
|  |  | Sometimes | 1781 (75) | 384 (76) | 353 (76) | 367 (74) | 337 (73) | 340 (76) |
|  |  | Never | 154 (6) | 44 (9) | 26 (6) | 34 (7) | 27 (6) | 23 (5) |

Abbreviations: BMI body mass index; CI confidence intervals; GCSE General Certificate of Secondary education; I:Cr urinary iodine to creatinine ratio; IQR interquartile range; SD standard deviation; UIC urinary iodine concentration. 1UK-equivalent qualifications were calculated from detailed overseas qualifications. GCSEs are standard assessments at age approx. 16. A-level assessments indicate education to approx. age 18. 2 Mean calculated from among those who were asked about diet in pregnancy (n=2776). 3 Gestational diabetes, hypertension or preeclampsia.

**Table 2 Child characteristics and cognitive outcomes according to maternal urinary iodine to creatinine ratio, n=7013 children**

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
|  | |  | **Maternal urinary iodine to creatinine ratio (fifths)** | | | | |
|  | | **All participants** | **1 – lower iodine** | **2** | **3** | **4** | **5 – higher iodine** |
| I:Cr (µg/g), median (range) | | 83 (1 to 2283) | 45 (1 to 54) | 64 (54 to 73) | 83 (73 to 95) | 111 (92 to 136) | 174 (136 to 2283) |
| n | | 7013 | 1401 | 1401 | 1401 | 1401 | 1401 |
| Sex (male), n (%) | | 3594 (51) | 688 (49) | 712 (51) | 731 (52) | 732 (52) | 731 (52) |
| Early years foundation stage [n=5383] | |  |  |  |  |  |  |
|  | Total points, all sub-domains (range 17 to 51), median (IQR)1 | 34 (28 to 37) | 34 (27 to 36) | 34 (28 to 36) | 34 (29 to 37) | 34 (28 to 37) | 34 (29 to 37) |
|  | At expected/exceeded level for all sub-domains, n (%) | 3031 (56) | 605 (55) | 604 (54) | 612 (58) | 614 (58) | 596 (58) |
| Phonics assessment in Year 1 | |  |  |  |  |  |  |
|  | Total score (range 0 to 40), median (IQR)1 [n=5468] | 37 (33 to 39) | 36 (33 to 38) | 37 (33 to 39) | 37 (33 to 39) | 37 (33 to 39) | 37 (34 to 39) |
|  | Achieved the standard (32/40), n (%) [n=5587] | 4424 (79) | 887 (78) | 894 (78) | 874 (80) | 903 (80) | 866 (80) |
| Key Stage 1, pre 2016 (achieved standard), n (%)3 [n=1317] | |  |  |  |  |  |  |
|  | English | 1032 (78) | 214 (74) | 203 (77) | 219 (83) | 194 (78) | 202 (80) |
|  | Writing | 888 (67) | 196 (68) | 177 (68) | 190 (72) | 156 (63) | 169 (67) |
|  | Maths | 1024 (78) | 220 (76) | 196 (75) | 217 (82) | 197 (79) | 194 (77) |
|  | Reading | 1038 (79) | 214 (74) | 205 (78) | 220 (83) | 195 (79) | 204 (81) |
|  | Science | 1148 (87) | 245 (84) | 230 (88) | 231 (88) | 219 (88) | 223 (88) |
| Key Stage 1, from 2016 (achieved standard), n (%)3 [n=4162] | |  |  |  |  |  |  |
|  | Writing | 2963 (71) | 591 (71) | 602 (70) | 587 (72) | 607 (71) | 576 (72) |
|  | Maths | 3190 (77) | 627 (75) | 658 (76) | 626 (77) | 645 (76) | 634 (79) |
|  | Reading | 3149 (76) | 618 (74) | 651 (75) | 625 (77) | 642 (75) | 613 (77) |
|  | Science | 3389 (81) | 662 (79) | 696 (81) | 680 (84) | 683 (80) | 668 (84) |
| Strengths & difficulties questionnaire [n=1019] | |  |  |  |  |  |  |
|  | Total difficulty score (range 0 to 40), median (IQR)2 | 5 (2 to 9) | 5 (2 to 9) | 4 (2 to 8) | 4 (2 to 8) | 4 (2 to 9) | 5 (2 to 9) |
|  | Close to average (0 to 11), n (%) | 864 (85) | 209 (80) | 182 (88) | 173 (86) | 156 (85) | 144 (86) |
|  | Slightly raised (12 to 15), n (%) | 108 (11) | 38 (15) | 14 (7) | 23 (11) | 19 (10) | 14 (8) |
|  | High or very high (16 to 40), n (%) | 47 (5) | 13 (5) | 10 (5) | 6 (3) | 9 (5) | 9 (5) |
| Letter identification score, mean (SD)1 [n=1421] | | 106·8 (12·9) | 105·5 (12·7) | 106·3 (13·0) | 107·7 (13·1) | 106·5 (12·1) | 108·4 (13·2) |
| British Picture Vocabulary Scale, mean (SD)1 [n=1439] | | 101·0 (16·3) | 99·2 (14·3) | 100·2 (14·9) | 101·0 (16·7) | 101·9 (19·1) | 103·5 (16·7) |
| CKAT scores [n=1426] | |  |  |  |  |  |  |
|  | Overall standardised score, mean (SD)1 | 99·5 (10·8) | 99·8 (10·6) | 99·2 (10·8) | 99·1 (10·5) | 99·7 (10·7) | 99·9 (11·3) |
|  | Tracking without guide: root mean square, median (IQR)2 | 14·1 (10·1 to 21·7) | 14·2 (10·0 to 22·3) | 14·0 (10·3 to 21·0) | 14·4 (10·4 to 23·6) | 13·3 (10·1 to 19·8) | 14·1 (9·7 to 21·2) |
|  | Tracking with guide line: root mean square, median (IQR)2 | 20·8 (12·7 to 36·1) | 20·4 (12·2 to 35·3) | 20·5 (12·4 to 34·3) | 22·8 (13·1 to 39·0) | 19·4 (13·3 to 37·4) | 20·9 (12·8 to 35·8) |
|  | Aiming: path length time (seconds), median (IQR)2 | 2·0 (1·7 to 2·3) | 2·0 (1·7 to 2·2) | 2·0 (1·7 to 2·3) | 1·9 (1·7 to 2·3) | 2·0 (1·8 to 2·3) | 1·9 (1·7 to 2·3) |
|  | Steering: path accuracy (mm), median (IQR)2 | 2·1 (1·7 to 2·7) | 2·1 (1·7 to 2·8) | 2·1 (1·7 to 2·9) | 2·0 (1·6 to 2·6) | 2·1 (1·7 to 2·7) | 2·1 (1·7 to 2·7) |

Abbreviations: CI confidence intervals; CKAT Clinical Kinematic Assessment Tool; IQR interquartile range; SD standard deviation.

1Higher scores represent better attainment or more advanced understanding; 2Higher scores represent more difficulty or slower and less accurate kinematic ability; 3 The KS1 standard before 2016 is defined as 'Working securely at level 2B or beyond’. From 2016 the standard is defined as ‘Working at the expected standard or beyond’.

**References**

1. SACN. SACN Statement on iodine and health: Public Health England, 2014.

2. Zimmermann MB. The importance of adequate iodine during pregnancy and infancy. *World Review of Nutrition and Dietetics* 2016; 115: 118-124.

3. Venkatesh Mannar MG. Making salt iodization truly universal by 2020. *IDD Newsletter* May 2014.

4. Roberts C, Steer T, Maplethorpe N, Cox L, Meadows S, Nicholson S, et al. National Diet and Nutrition Survey. Results from years 7 and 8 (combined) of the rolling programme (2014/2015 to 2015/2016). A survey carried out on behalf of Public Health England and the Food Standards Agency. Available online <https://assets.publishing.service.gov.uk/government/uploads/system/uploads/attachment_data/file/699241/NDNS_results_years_7_and_8.pdf> [Accessed December 2019] London: Public Health England; 2018.

5. Andersson M, de Benoist B, Delange F, Zupan J. Prevention and control of iodine deficiency in pregnant and lactating women and in children less than 2-years-old: conclusions and recommendations of the Technical Consultation. *Public Health Nutrition* 2007; 10(12a): 1606-1611.

6. van Mil NH, Tiemeier H, Bongers-Schokking JJ, Ghassabian A, Hofman A, Hooijkaas H, et al. Low urinary iodine excretion during early pregnancy is associated with alterations in executive functioning in children. *Journal of Nutrition* 2012; 142(12): 2167-2174.

7. Bath SC, Steer CD, Golding J, Emmett P, Rayman MP. Effect of inadequate iodine status in UK pregnant women on cognitive outcomes in their children: results from the Avon Longitudinal Study of Parents and Children (ALSPAC). *Lancet* 2013; 382(9889): 331-337.

8. Ghassabian A, Steenweg-de Graaff J, Peeters RP, Ross, HA, Jaddoe VW, Hofman A, et al. Maternal urinary iodine concentration in pregnancy and children's cognition: results from a population-based birth cohort in an iodine-sufficient area. *BMJ Open* 2014; 4(6): e005520.

9. Hynes KL, Otahal P, Burgess JR, Oddy WH, Hay I. Reduced educational outcomes persist into adolescence following mild Iodine deficiency in utero, despite adequacy in childhood: 15-Year follow-up of the gestational iodine cohort investigating auditory processing speed and working Memory. *Nutrients* 2017; 9(12): 1354.

10. Murcia M, Espada M, Julvez J, Llop S, Lopez-Espinosa MJ, Vioque J, et al. Iodine intake from supplements and diet during pregnancy and child cognitive and motor development: the INMA Mother and Child Cohort Study. *Journal of Epidemiology and Community Health* 2018; 72(3): 216-222.

11. Markhus MW, Dahl L, Moe V, Abel HM, Brantsaeter LA, Oyen J, et al. Maternal iodine status is associated with offspring language skills in infancy and toddlerhood. *Nutrients* 2018; 10(9): 1270.

12. Robinson SM, Crozier SR, Miles EA, Gale CR, Calder PC, Cooper C, et al. Preconception maternal iodine status is positively associated with IQ but not with measures of executive function in childhood. *Journal of Nutrition* 2018; 148(6): 959-966.

13. Zhou SJ, Condo D, Ryan P, Howell S, Skeaff SA, Anderson PJ, et al. Association between maternal iodine intake in pregnancy and childhood neurodevelopment at age 18 months. *American Journal of Epidemiology* 2019; 188(2): 332-338.

14. Levie D, Korevaar TIM, Bath SC, Murcia M, Dineva M, Llop S, et al. Association of maternal iodine status with child IQ: a meta-analysis of individual-participant data. *Journal of Clinical Endocrinology and Metabolism* 2019; 104(12): 5957-5967.

15. Gowachirapant S, Jaiswal N, Melse-Boonstra A, Galetti V, Stinca S, Mackenzie I, et al. Effect of iodine supplementation in pregnant women on child neurodevelopment: a randomised, double-blind, placebo-controlled trial. *Lancet Diabetes and Endocrinology* 2017; 5(11): 853-863.

16. Wright J, Small N, Raynor P, Tuffnell D, Bhopal R, Cameron N, et al. Cohort Profile: the Born in Bradford multi-ethnic family cohort study. *International Journal of Epidemiology* 2013; 42(4): 978-991.

17. Caldwell KL, Makhmudov A, Jones RL, Hollowell JG. EQUIP: A worldwide program to ensure the quality of urinary iodine procedures. *Accreditation and Quality Assurance* 2005; 10(7): 356-361.

18. Li C, Peng S, Zhang X, Xie X, Wang D, Mao J, et al. The urine iodine to creatinine as an optimal index of iodine during pregnancy in an iodine adequate area in China. *Journal of Clinical Endocrinology and Metabolism* 2016; 101(3): 1290-1298.

19. Born in Bradford. Born in Bradford data dictionary. Starting school: all measures and administrative information <https://borninbradford.nhs.uk/wp-content/uploads/Starting_School_StartingSchool_Full_Dict.pdf>. [Accessed December 2019] Bradford: Bradford Institute for Health Research, 2018.

20. Cotzias E, Whitehorn T, STA Teacher Assessment & Moderation team. Topic Note: Results of the Early Years Foundation Stage Profile (EYFSP) pilot. Research report. Available online <https://assets.publishing.service.gov.uk/government/uploads/system/uploads/attachment_data/file/190639/DFE-RR291.pdf>. [Accessed December 2019], 2013.

21. Shire K, Andrews E, Barber S, Bruce A, Corkett J, Hill LJB, et al. Starting School: a large-scale start of school assessment within the ‘Born in Bradford’ longitudinal cohort [version 1; peer review: 1 approved, 1 approved with reservations]. *Wellcome Open Research* 2020; 5:47.

22. Goodman R. The Strengths and Difficulties Questionnaire: a research note. *Journal of Child Psychology and Psychiatry and Allied Disciplines* 1997; 38(5): 581-586.

23. Knudsen N, Christiansen E, Brandt-Christensen M, Nygaard B, Perrild H. Age- and sex-adjusted iodine/creatinine ratio. A new standard in epidemiological surveys? Evaluation of three different estimates of iodine excretion based on casual urine samples and comparison to 24 h values. *European Journal of Clinical Nutrition* 2000; 54(4): 361-363.

24. Rogers WH. Regression standard errors in clustered samples. *Stata Technical Bulletin* 1993; 13: 19-23.

25. Harrell FE. Regression modeling strategies with applications to linear models, logistic regression, and survival analysis. New York: Springer; 2001.

26. WHO. Assessment of iodine deficiency disorders and monitoring their elimination. A guide for programme managers. Available online <https://apps.who.int/iris/handle/10665/43781>. [Accessed June 2020] Third ed. France: World Health Organization, 2007.

27. Konig F, Andersson M, Hotz K, Aeberli I, Zimmermann MB. Ten repeat collections for urinary iodine from spot samples or 24-hour samples are needed to reliably estimate individual iodine status in women. *Journal of Nutrition* 2011; 141(11): 2049-2054.

28. Perrine CG, Cogswell ME, Swanson CA, Sullivan KM, Chen TC, Carriquiry AL, et al. Comparison of population iodine estimates from 24-hour urine and timed-spot urine samples. *Thyroid* 2014; 24(4): 748-757.

29. Velasco I, Bath SC, Rayman MP. Iodine as essential nutrient during the first 1000 days of life. *Nutrients* 2018; 10(3): 290.

30. Moss G, Washbrook L. Understanding the gender gap in literacy and language development. Bristol Working Papers in Education #01/2016. Available online <http://www.bristol.ac.uk/media-library/sites/education/documents/bristol-working-papers-in-education/Understanding%20the%20Gender%20Gap%20working%20paper.pdf> [Accessed November 2018], 2016.

31. Bath SC, Furmidge-Owen VL, Redman WG, Rayman MP. Gestational changes in iodine status in a cohort study of pregnant women from the United Kingdom: season as an effect modifier. *American Journal of Clinical Nutrition* 2015; 101:1180-1187.

32. Costeira MJ, Oliveira P, Ares S, de Escobar GM, Palha JA. Iodine status of pregnant women and their progeny in the Minho region of Portugal. *Thyroid* 2009; 19(2):157-163.

**Supporting information**

eFigure 1: Flow chart of participant inclusions and exclusions

eTable 1: Urinary iodine status of healthy pregnant UK populations – all report iodine insufficiency according to World Health Organization-outlined thresholds

eTable 2: Details of socioeconomic status and maternal education levels

Supplementary Methods

eTable 3: Missing data imputed

eFigure 2: A directed acyclic graph used to identify confounders and competing exposures in the association between maternal iodine concentration and child cognitive outcomes

eTable 4: Characteristics of mothers with usable urine samples and the rest of cohort

eTable 5: Maternal characteristics according to ethnic group, n= 6971 mothers

eTable 6: Predicted estimates (continuous outcomes) and percent at the threshold (binary outcomes) (99% CIs) at the 25th, 50th and 75th centiles of iodine concentration for 'typical' participants

eFigure 3: Associations between maternal I:Cr and the estimated EYFS score in the full sample of participants and subgroups

eFigure 4: Associations between maternal I:Cr and the estimated Phonics score in the full sample of participants and subgroups

eFigure 5: Associations between maternal I:Cr and the estimated percent achieving the EYFS standards in key domains in the full sample of participants

eFigure 6: Associations between maternal I:Cr and the estimated percent achieving the KS1 reading standards in pre-2016 assessments in the full sample and subgroups

eFigure 7: Associations between maternal I:Cr and SDQ total, internalising and externalising scores and the estimated percent being scored as at least slightly raised in the full sample

eFigure 8: Associations between maternal I:Cr and the estimated percent being graded as at least slightly raised with the SDQ in the full sample and subgroups

eFigure 9: Associations between maternal I:Cr and BPVS, Letter identification and CKAT outcome measures in the full sample of participants