Birthweight associations with parental obesity: retrospective analysis of 1,778 singleton term births following assisted reproductive treatment

Nicole O. McPherson, Ph.D., Andrew D. Vincent, Ph.D., Deirdre Zander-Fox, Ph.D., and Jessica A. Grieger, Ph.D.

Objective: To determine the association of combined parental preconception overweight and obesity on infant birthweight.

Design: Retrospective study of fresh in vitro fertilization or intracytoplasmic sperm injection cycles (2009–2017).

Setting: Repromed, South Australia, assisted reproductive technology clinic.

Patients: Couples undergoing in vitro fertilization/intracytoplasmic sperm injection insemination with their own gametes and transfer of a fresh single blastocyst (N = 1,778).

Intervention(s): None.

Main Outcome Measures: Parental body mass index (BMI) was recorded prior to cycle initiation. Infant birthweight was recorded at delivery. The impact of parental obesity and their interaction on first singleton term (≥ 37 weeks’ gestation) birthweight was assessed using linear regressions assessing nonlinearity and a pairwise linear interactions.

Results: In the base model where parental BMI is assumed linear, there was strong evidence for higher birthweight with increasing maternal BMI (11.2 g per maternal kg/m²; 95% confidence interval, 7.2, 15.1) but not paternal BMI. The inclusion of a pairwise linear interaction indicated that paternal BMI attenuates the positive association between maternal BMI and infant birthweight (interaction /C0 0.88; 95% confidence interval, /C0 1.49, /C0 0.27). The inclusion of nonlinear maternal BMI terms did not change the conclusions.

Conclusions: Increases in the mean infant birthweight associated with maternal obesity are attenuated when the father is obese. While maternal BMI contributed more to the mean infant birthweight than paternal BMI, a couple-centered approach to preconception health advice is recommended, given the documented relationships between parental obesity and childhood weight beyond infancy. Further studies in both assisted reproductive technology and general population cohorts assessing the parental BMI interaction on infant birthweight are warranted. (Fertil Steril Rep® 2021;2:405–12. ©2021 by American Society for Reproductive Medicine.)

Key Words: Maternal obesity, paternal obesity, offspring birthweight, body mass index, preconception

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Obesity is a significant public health concern. There is a rising trend for increased body mass index (BMI) across all age groups, with obesity rates tripling over the past 40 years (1). A global systematic analysis found that men and women have gained weight across all age groups, with most rapid gains occurring during prime reproductive years (ages 20 and 40 years) (2). In 2014, 38.9 million pregnant women were estimated to be overweight, and 14.6 million were estimated to be obese (3). In Australia, approximately 50% of women and 70% of men who contributed to a pregnancy in 2017 were overweight or obese (4).

It is established that maternal prepregnancy BMI has a significant impact on infant birthweight, and this is irrespective of whether couples conceived naturally or through assisted reproductive technology (ART). In non-ART populations, maternal overweight or obesity increases the likelihood of an infant being born large for gestational age (odds ratio [OR], 1.45; 95% confidence interval [CI], 1.29, 1.63; and OR, 1.88; 95% CI, 1.67, 2.11, respectively) or with macrosomia (OR, 1.70; 95% CI, 1.55, 1.87; and OR, 2.92; 95% CI, 2.67, 3.20, respectively) (5). In comparison, the risk of delivering a small-for-gestational-age baby increases in mothers who are underweight (OR, 1.67; 95% CI, 1.49, 1.87) but decreases in those who are overweight (OR, 0.71; 95% CI, 0.66, 0.76) or obese mothers (OR, 0.88; 95% CI, 0.78, 0.99) (5). In women requiring ART, both maternal underweight and obesity are associated with an increased risk of low birthweight (adjusted risk ratio, 1.39; 95% CI, 1.25, 1.54; and adjusted risk ratio, 1.26; 95% CI, 1.20, 1.33, respectively) (6). The variance in results between studies is likely due to the differences seen between natural conception and ART cohorts, with several meta-analyses and large cohort studies showing that the mean birthweight in ART singletons is lower than in naturally conceived singletons (7, 8).

However, the potential impact of paternal BMI is rarely considered in these studies, despite a small body of evidence suggesting that paternal preconception overweight and obesity may also contribute to infant birthweight (9, 10). Furthermore, evidence for an interaction between the preconception obesity status of parents and infant birthweight is not possible to discern from existing studies, given that obesity tends to aggregate in family units so that obese mothers commonly have obese partners (11). One study including 2,980 parent–offspring trios reported that maternal BMI was a stronger predictor of ponderal index and birthweight than paternal BMI (12). A limitation of this study was that parental preconception body weight was estimated retrospectively by the mother at the time of child data collection at 3–18 years (12).

Therefore, while it is clearly evident that maternal preconception BMI affects infant birthweight, the influence of paternal preconception BMI remains unclear. Furthermore, it is unknown whether there is an additional effect on infant birthweight if both parents are overweight or obese. We hypothesized that the combination of both maternal and paternal preconception overweight/obesity has a greater impact on infant birthweight than either independent parental effect. This study aimed to assess the independent and combined effects of maternal and paternal preconception overweight and obesity on the mean infant birthweight utilizing an ART cohort where preconception parental BMI is routinely collected.

MATERIALS AND METHODS

Human Ethics

The Scientific Advisory Committee of Repromed (Adelaide, Australia) approved the retrospective study (November 14, 2019). The study was exempt from Human Research Ethics Committee review at the University of Adelaide. Patients had previously given written consent to allow their records to be accessed for low-risk retrospective investigation.

Study Population and Data Collection

The study involved the retrospective analysis of fresh cycles from 2009 to 2017 at Repromed (Dulwich, South Australia, and Darwin, Northern Territory, clinics). Cycles including in vitro fertilization (IVF) or intracytoplasmic sperm injection (ICSI) with autologous sperm and eggs and the transfer of a single blastocyst embryo were assessed (Supplemental Fig. 1, available online). First singleton term births (≥37 weeks’ gestation) with birthweights recorded were included in the analysis. Preterm births (<37 weeks’ gestation), twin births, and second pregnancies from the same patient couple were excluded from the analysis (Supplemental Fig. 1). Parental, demographic, and treatment factors that are known to influence birthweight including maternal (13) and paternal age (14), socioeconomic status (15) (assessed using the Socio-Economic Indexes for Areas [SEIFA]) (16), insemination method (17), sex of baby (18), delivery method (19), and gestational age (20) were collected from case notes. A high SEIFA score indicates greater social advantage, while a low score indicates relatively greater disadvantage; the average SEIFA score is 1,000, and the middle two-thirds of the SEIFA scores generally fall between approximately 900 and 1,100 (16). Infertility diagnosis was also collected from case notes; however, it was not included in the final modeling analysis as it has been previously shown to not be associated with birthweight in term pregnancies (21, 22).

All data including infant birthweight (g), gestational age (weeks), sex (male/female), twin deliveries, and delivery method (vaginal/cesarean) were supplied by the treating obstetrician as per the ART treatment act that indicates mandatory reporting to the Australian and New Zealand Assisted Reproduction Database.

Assessment of Parental BMI

Body mass index of both parents was recorded before cycle initiation as part of routine clinical practice at Repromed. Both maternal and paternal heights were measured with a stadiometer (cm), and weight (kg) was measured with electronic scales. Body mass index was calculated using the formula weight/height² and categorized based on the World Health Organisation criteria as underweight (<18.5 kg/m²), normal weight (18.5–24.9 kg/m²), overweight (25.0–29.9 kg/m²), and obesity (>30.0 kg/m²). Obesity was subclassified as
obesity class I (30.0–34.9 kg/m²), obesity class II (35.0–39.9 kg/m²), and obesity class III (>40 kg/m²).

**IVF Protocol**

Women underwent a gonadotropin-releasing hormone antagonist protocol of treatment with vaginal progesterone gel (Crinone)/estradiol valerate luteal support or human-derived human chorionic gonadotropin luteal support (Premyn), as previously described (23). Over the course of the study (2009–2017), there were no substantive changes to laboratory protocols, including in culture media, consumables, gas phase, or equipment used including incubator type. During this time frame, the clinic was relatively static in its clinical protocols and policies. Extended culture was standard protocol (cleavage-stage transfer was rarely utilized), and stimulation regimes were also relatively static (antagonist protocol was standard). Eggs were fertilized by either standard IVF or ICSI in fertilization medium (G-IVF-PLUS, Vitrolife, Gothenburg, Sweden). Embryos were cultured using the sequential culture media system supplied by Vitrolife at 6% CO₂, 5% O₂, and 89% N₂, wherein cleavage-stage embryos were grown until day 3 in G1 PLUS and then moved into G2 PLUS, which supported blastocyst development until embryo transfer on day 4 or 5. The best morphologically graded blastocyst was transferred using EmbryoGlue transfer medium (Vitrolife). Patients were in the care of their treating IVF physician until confirmation of a viable pregnancy following ultrasound at 6–8 weeks’ gestation and then referred on to primary obstetric care.

**Statistical Methods**

For continuous demographic, treatment, and outcome factors, means (standard deviations) and medians (ranges) are reported, and for discrete factors, frequencies (percentages) are reported. Associations between paternal and maternal BMI and infant birthweight were assessed using linear regressions, adjusting for baby sex (male or female), gestational age, delivery method (cesarean or vaginal), transfer method (IVF or ICSI), year of birth, maternal age, paternal age, and parental SEIFA score. Nonlinear associations were modeled using restricted cubic splines (knots at 5th, 35th, 65th, and 95th percentiles) for gestational age. This base model (M0) was extended in the following three stages: model M1, initially with the inclusion of the pairwise linear interaction of parental BMIs; model M2, then with the inclusion of nonlinear terms for maternal BMI; and model M3, finally with the inclusion of nonlinear terms for paternal BMI, that is, this final model included nonlinear terms for both parental BMIs and their pairwise linear interaction. The nonlinear associations were also modeled using restricted cubic splines with four knots. Multiple imputation using chained equations (100 datasets were imputed each with 100 iterations) was employed to account for the substantial missing parental BMI data (18% and 33% for maternal and paternal data, respectively). Analyses were performed in R (version 3.6.3) using the mice and rms packages. A P value of <.05 was considered statistically significant.

**RESULTS**

**Patient Demographics**

A total of 1,778 couples were included in the analysis (Supplemental Fig. 1). The median age of mothers was 32 (range, 20.0–45.0) years, which was lower than that of fathers (35 years; range, 20.0–65.0) (Table 1). The median BMI of mothers (24.4 kg/m²; range, 16.2–55.9) was in the normal weight category, but BMI spanned from underweight (<18.5 kg/m²) to obese class III (>40 kg/m²) categories. The median BMI of fathers (27.4 kg/m²; range, 17.3–54.2) was in the overweight category and also spanned from underweight to obese class III. Moreover, 33% of paternal BMI data was missing compared with 18% of maternal BMI data. The mean SEIFA score (mean, 994; standard deviation, 72) was slightly lower than the Australian benchmark of 1,000, reflecting social disadvantage in this cohort. Male factor infertility was the most common contributing infertility diagnosis (53%) in the cohort. Intracytoplasmic sperm injection insemination was used in over 80% of cases with delivery method (vaginal vs. cesarean) and infant sex (female vs. male) split approximately 50%. Table 1 presents demographic summary statistics, with Supplemental Table 1 (available online) presenting these statistics by parental obesity categories.

**Parental Preconception BMI and Infant Birthweight**

In the base model of infant birthweight (model M0) in which the influence of parental BMI was assumed linear, there was strong evidence for higher birthweight with increasing maternal BMI, with a mean increase of 11.2 g per maternal kg/m² (95% CI, 7.2, 15.1; P < .001; Table 2). In this base model, there was no evidence of an association between paternal BMI and birthweight (P = .44, Table 2). The inclusion of a pairwise linear interaction (model M1), however, indicated that paternal BMI appears to attenuate the positive association between maternal BMI and infant birthweight (interaction; 95% CI, −1.49, −0.27; P = .005; Table 2). As shown in Figure 1A, Supplemental Fig. 2A and Table 3, the estimated mean (95% CI) birthweights only increased from 3.33 kg (3.25, 3.42) to 3.43 kg (3.36, 3.50) in mothers who had BMIs equal to 20 kg/m² as compared with 35 kg/m², when the father was obese (BMI = 35 kg/m²). Comparatively, with a paternal BMI of 20 kg/m², birthweight increased from 3.19 kg (3.11, 3.27) to 3.48 kg (3.38, 3.58) with increasing maternal BMI (20 vs. 35 kg/m²). The inclusion of nonlinear maternal BMI terms (model M2) did not qualitatively change these conclusions (paternal linear interaction P = .02, Fig. 1B, Supplemental Fig. 2B and Table 2). However, the interaction did become nonsignificant (P = .15) when both parental BMIs were modeled nonlinearly (model M3). This addition of nonlinear paternal BMI added negligible explanatory value (P = .41, Table 2), with no qualitative difference in the estimated mean (95% CI) birthweights for normal–weight versus obese parents. For example, in this full model, as presented in Figure 1C, Supplemental Fig. 2C and Table 3, the estimated birthweights increased from 3.18 kg (3.07, 3.29) to 3.43 kg (3.29, 3.57) for an increase in mothers’ BMI from 20 to 35 kg/m², when the father had
TABLE 1
Summary of parental demographics, treatment choices, and birth outcomes.

| Parental characteristics | N = 1,778 |
|--------------------------|-----------|
| **Age and BMI**          |           |
| Maternal age (years)     | Median (range) 33.0 (20.0, 45.0) |
|                         | Mean (SD) 32.8 (4.12) |
| Paternal age (years)     | Median (range) 35.0 (20.0, 65.0) |
|                         | Mean (SD) 36.1 (6.3) |
| **Maternal BMI (kg/m²)** | Median (range) 24.4 (16.2, 55.9) |
|                         | Mean (SD) 25.9 (6.0) |
| Missing                  | 315 (18%) |
| <18.5 kg/m²              | 40 (2%) |
| 18.5–24.9 kg/m²          | 754 (42%) |
| 25–29.9 kg/m²            | 371 (21%) |
| 30–34.9 kg/m²            | 155 (9%) |
| 35–39.9 kg/m²            | 91 (5%) |
| >40 kg/m²                | 52 (3%) |
| **Paternal BMI (kg/m²)** | Median (range) 27.4 (17.3, 54.2) |
|                         | Mean (SD) 28.1 (4.6) |
| Missing                  | 585 (33%) |
| <18.5 kg/m²              | 3 (1%) |
| 18.5–24.9 kg/m²          | 283 (16%) |
| 25–29.9 kg/m²            | 572 (32%) |
| 30–34.9 kg/m²            | 238 (13%) |
| 35–39.9 kg/m²            | 68 (4%) |
| >40 kg/m²                | 29 (2%) |
| **Couple BMI (kg/m²)**   | Both <30 kg/m² 747 (42%) |
|                         | Maternal <30 kg/m² 207 (12%) |
|                         | paternal >30 kg/m² 117 (7%) |
|                         | Maternal >30 kg/m² 214 (12%) |
|                         | paternal missing 62 (3%) |
|                         | Maternal >30 kg/m² 2 (1%) |
|                         | Maternal >30 kg/m² 4 (1%) |
|                         | Both missing 309 (17%) |
| **SEIFA**                | Median (range) 1,000 (673, 1163) |
|                         | Mean (SD) 994 (72) |
|                         | Missing 35 (2%) |
| **Infertility diagnosis**| Tubal factor 147 (8%) |
|                         | Endometrial factor 121 (7%) |
|                         | Male factor 949 (53%) |
|                         | Other 590 (33%) |
|                         | Unexplained 380 (21%) |
| **Birth factors**        | Insemination method IVF 278 (16%) |
|                         | ICSI 1,500 (84%) |
|                         | Delivery method Vaginal 1,041 (59%) |
|                         | Cesarean 734 (41%) |
|                         | Missing 3 (1%) |
|                         | Gestational length (weeks) Median (range) 39.1 (37.0, 42.1) |
|                         | Mean (SD) 39.2 (1.1) |

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**DISCUSSION**

In a retrospective study of 1,778 singleton term births following ART, we were unable to detect increasing birthweight when both parents were obese, as compared with just one parent alone. That is, while infants are heavier when born to overweight or obese mothers, this association appears to be substantially attenuated in the presence of paternal obesity. Importantly, the estimated increase in birthweight with maternal obesity (with normal paternal BMI) was double than that observed with paternal obesity (and normal maternal BMI), demonstrating a much larger contribution of maternal BMI than that of paternal BMI to infant birthweight.

To our knowledge, this is the first study assessing the possibility of a combined contribution of maternal and paternal preconception BMIs on infant birthweight. The strengths of our study include the use of a database in which preconception health, IVF cycle outcomes, and pregnancy rates were registered prospectively, thereby minimizing selection bias. Body mass index was calculated preconception from clinically recorded measurements of maternal and paternal weights and heights; the analysis only included first singleton term births; and the large population size from a singular ART unit limited variability in clinical protocols. The limitations of our study include the retrospective study design, which limits the ability to control and collect some key parental factors that can influence infant birthweight, including parental smoking, ethnicity, or infertility duration (24–27). The use of a subfertile cohort means that our results may only be generalizable to couples undergoing ART, although infertility diagnosis has previously been shown to normal weight (BMI = 20 kg/m²). This is twice the change in birthweights that was observed for when fathers were obese (BMI = 35 kg/m²), for which the increase in birthweight was 3.30 kg (3.20, 3.40) to 3.42 kg (3.34, 3.50) for the same change in mothers’ BMI. The estimates for the four models, including the estimates for the adjustments (i.e., parental age, SEIFA score, baby sex, gestational age, delivery method, transfer method, and year of delivery), are shown in Supplemental Table 2 (available online).
minimally influence infant birthweight in term pregnancies (21, 22). There was no prior power calculation performed for this analysis, although it is known that large sample sizes are required to reliably detect interactions.

Contrary to our hypothesis, we found no additional effect of having two overweight or obese parents on infant birthweight beyond the effect of maternal obesity. Evidence from our rodent model of obesity also indicates that the effect of parental obesity on infant birthweight is unlikely to be additive but is instead a combination of both maternal and paternal phenotypes (28). This is evident in the current dataset where the effect of maternal obesity on infant birthweight was substantially attenuated in the presence of paternal obesity. This is because infants born to obese fathers start out heavier (approximately 150 g). While we found no additional effect of combined parental BMI on infant birthweight, additive effects may become evident as infants grow. For instance, the Raine cohort in Western Australia demonstrated that parental obesity was the strongest predictor of offspring adult BMI (29), while the Midspan Family Study in Scotland...

### TABLE 2

The association of parental body mass index with infant birthweight, examining the influence of nonlinearity and a pairwise linear interaction.

| Parental interactions | M0 (linear BMI) | M1 (linear + int) | M2 (non-lin mat-BMI + int) | M3 (non-lin BMI + int) |
|-----------------------|-----------------|-------------------|---------------------------|-----------------------|
|                       | Est (g) [95% CI] | P value           | Est (g) [95% CI]          | P value               |
| Mat BMI (linear)      | 11.2 [7.2, 15.1] | < .001            | 12.2 [8.2, 16.2]          | < .001                |
| Pat BMI (linear)      | 2.0 [−3.1, 7.1]  | .44               | 4.4 [−1.0, 9.9]           | .13                   |

Note: BMI = body mass index; Mat = maternal; Pat = paternal; lin = linear; int = interaction. *A single P value is reported for all nonlinear terms. The base model M0 includes just the two parental BMIs linearly. Model M1 extends M0 with the inclusion of the pairwise linear interaction of parental BMIs. Model M2 extends M1 with the inclusion of nonlinear terms for maternal BMI. Model M3 is the full model with nonlinear terms for both parental BMIs and their pairwise linear interaction. The following adjustment factors were included in the models: gestational age, parental age, Socio-Economic Indexes for Areas score, year of delivery, and delivery method.

**FIGURE 1**

Estimated mean infant birthweight by maternal body mass index (BMI). The interaction is illustrated by varying maternal BMI with paternal (Pat) BMI set at 20 (blue) and 35 (red) kg/m², respectively. (A) Model M1 with both linear paternal BMIs, (B) model M2 with nonlinear maternal BMI, and (C) model M3 with both nonlinear paternal BMIs. Gray circles are observed birthweights. Solid lines are the estimated means, and dashed lines are the 95% confidence intervals. Presented mean estimates are for covariates: female baby; gestational age, 3 weeks; vaginal birth in 2012; in vitro fertilization insemination; maternal age, 33 years; paternal age, 35 years; and Socio-Economic Indexes for Areas score, 1000.

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revealed that adult offspring from two obese parents had a higher risk of cardiovascular diseases compared with offspring without obese parents (30). In contrast, a study assessing 2,980 parent–offspring trios showed that the effect of parental preconception BMI on child and adolescent BMI was minimal (12) and suggested that while parental obesity may play a role in infant birthweight, family environment likely plays a much larger role in cardio–metabolic disease risk in adult life. The clinical relevance of our results are unclear (i.e., approximately 290-g increase in birthweight is unlikely to impact clinical outcomes); however, birthweight is reported to play a significant role in the establishment of adolescent and early adulthood BMI (29, 31). Therefore, if obesity aggregates within families, then a focus on preconception planning for “healthy couples” and emphasizing the need to improve lifestyle for the family unit prior to pregnancy is recommended. This may help shape and establish later life habits to support long-term health of future generations.

There is a large body of literature demonstrating the impact of maternal BMI on infant birthweight (32), and there is some evidence that paternal BMI also impacts infant birthweight (9, 10). Unfortunately, much of the literature on paternal BMI includes self-reported paternal height and weight from the mother during pregnancy, at birth, or when the child was a toddler, rather than preconception, with adjustments for potential confounders often inadequate (33, 34). Thus, the impact of paternal BMI remains unclear. Some (35, 36) but not all (37–39) studies have demonstrated an association between paternal BMI and infant birthweight, with similar mixed reports seen on the extreme ends of infant birthweight (small for gestational age or large for gestational age) (22, 40, 41). In our study, we found that paternal BMI only minimally impacted infant birthweight, with a paternal effect only evident when paternal BMI was added to the regression models as a pairwise linear interaction (4.43 g for every 1-unit increase in paternal BMI). The lack of consensus in the reported effects of paternal overweight and obesity on infant birthweight highlights the necessity for further large cohort studies in both ART and naturally conceived populations, ensuring correct clinical measurements of preconception paternal BMI.

The mechanism on how parental obesity is altering infant birthweight is likely due to a combination of genetic and epigenetic factors (noncoding ribonucleic acid (RNAs) and deoxyribonucleic acid (DNA) and histone methylation) delivered by sperm and eggs at fertilization (42–44) and the relationship between in utero fetal programming by nutritional stimuli (45, 46). A number of genes are known to play a part in the heritability of weight (47, 48); however, these genetic loci do not fully account for the transmission, indicating that programming effects go beyond underlining genetics. Maternal obesity during gestation has been associated with an altered expression of a number of circulating microRNAs with levels directly correlating with changes in placental weight and infant birthweight (49), while a number of studies in animal models and humans directly show a link between paternal obesity at conception, sperm epigenetic changes, and altered fetal phenotypes (36, 50–54). These studies collectively indicate that the parental programming effect to infant birthweight goes beyond that of a shared living environment, with preconception factors able to influence the health of subsequent children.

In conclusion, utilizing close to 1,800 singleton term births from an Australian ART cohort, our results demonstrate that increases in mean infant birthweight associated with maternal obesity are attenuated when the father is obese. While maternal BMI contributed more to the mean infant birthweight than paternal BMI, a couple–centered approach to preconception health advice could be valuable, given the relationships documented between parental obesity and childhood weight beyond infancy. Further studies in both ART and general population cohorts assessing the interaction of maternal and paternal preconception BMIs on infant birthweight to support or refute our findings are warranted.

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| Maternal BMI (kg/m²) | Paternal BMI (kg/m²) | M1: birthweight (g) [95% CI] | M2: birthweight (g) [95% CI] | M3: birthweight (g) [95% CI] |
|---------------------|---------------------|-----------------------------|-----------------------------|-----------------------------|
| 20                  | 20                  | 3191 [3112, 3270]           | 3176 [3095, 3256]           | 3176 [3066, 3286]           |
| 35                  | 20                  | 3481 [3382, 3580]           | 3457 [3354, 3560]           | 3427 [3287, 3567]           |
| 20                  | 35                  | 3335 [3250, 3421]           | 3309 [3215, 3403]           | 3296 [3196, 3395]           |
| 35                  | 35                  | 3427 [3355, 3500]           | 3412 [3336, 3487]           | 3422 [3344, 3499]           |

Note: BMI = body mass index. Model M1 extends M0 with the inclusion of the pairwise linear interaction of parental BMIs. Model M2 extends M1 with the inclusion of nonlinear terms for maternal BMI. Model M3 is the full model with both nonlinear terms for both parental BMIs and their pairwise linear interaction. The following adjustment factors were included in the models: gestational age, parental age, Socio–Economic Indexes for Areas score, year of delivery, and delivery method.

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