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Title Page

Gestational Dyslipidaemia and the Risk of Extreme Birth Weight: A

3 Systematic Review and Meta-analysis

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20 Abstract

Background

- Low and high birthweight is known to increase the risk of acute and longer term adverse
- outcomes, such as stillbirth, infant mortality, obesity, type 2 diabetes, and cardiovascular
- diseases. Gestational dyslipidaemia is associated with a numbers of adverse birth outcomes,
- but evidence regarding on birth weight is still inconsistent to reliably inform clinical practice
- and treatment recommendations.

Objective

- 28 To explore the relationship between maternal gestational dyslipidaemia and neonatal health
- 29 outcomes namely, birth weight, metabolic factors, and inflammatory parameters.

30 Methods

- We searched systematically Embase, MEDLINE, PubMed, CINAHL Plus, and Cochrane
- Library up to 1st August 2016 (with an updated search in MEDLINE at the end of July 2017),
- 33 for longitudinal studies that assessed the association of maternal lipid levels during
- 34 pregnancy with neonatal birth weight, or metabolic and inflammatory parameters up to 3
- years old.

36 Results

- 37 Data from 46 publications including 31,402 pregnancies suggests that maternal high
- 38 triglycerides and low high-density-lipoprotein cholesterol levels throughout pregnancy are
- 39 associated with increased birth weight, higher risk of large-for-gestational age and
- 40 macrosomia; and lower risk of small-for-gestational age. The findings were consistent across
- 41 the studied populations, but stronger associations were observed in women who were
- 42 overweight or obese prior to pregnancy.

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- LBW: low birth weight
- SGA: small for gestational age
- LGA: large for gestational age
- GDM: gestational diabetes mellitus
- **RCT**: Randomised Controlled Trial

- total cholesterol

 DL: high-density lipoprotein

 DL: low-density lipoprotein

 VLDL: very low-density lipoprotein

 *riglycerides

- IL-6: interleukin 6
- TNF-α: Tumour Necrosis Factor alpha
- 11β HSD1: 11-beta-Hydroxysteroid Dehydrogenase Type 1
- CRP: C-reactive protein
- T1: the first trimester
- T2: the second trimester

- 69 T3: the third trimester
- 70 mg/dL: milligrams per decilitre
- 71 mmol/L: millimoles per litre
- 72 RC: regression coefficients
- 73 OR: odds ratio
- 74 MD: mean difference
- 75 GWAS: genome-wide association study

Introduction

Low and high birth weight has been linked to the risk of stillbirth and infant mortality. In a longer life course, both low birth weight(LBW) or small for gestational age(SGA), and large for gestational age(LGA) or macrosomia are known to increase the future risk of obesity, type 2 diabetes, and cardiovascular disease.^{2, 3} The estimated prevalence of macrosomia in developed countries varies from 5% to 20%, and a parallel increase in macrosomic births was observed in both developed and developing countries over the last two to three decades.⁴ These life course associations have often been attributed to the impact of an adverse intrauterine environment, particularly, fuels (glucose, lipids, and amino acids) transported from the maternal end.⁵ Previous reviews have shown that maternal obesity and gestational diabetes mellitus(GDM) are two identified risk factors of low and high birthweight.⁶⁻⁸ However, as one of common metabolic disorders, the adverse effects of gestational dyslipidaemia on neonates birth weight/birth weight centiles are not widely recognized in clinical practice. Dyslipidaemia has been considered a risk factor for a number of adverse health outcomes, in particular cardiovascular disease and type 2 diabetes.^{9, 10} Previous reviews have shown that dyslipidaemia during pregnancy are associated with increased risk of GDM, preeclampsia, and pre-term delivery¹¹⁻¹³, but epidemiological evidence on birthweight is conflicting¹⁴⁻¹⁶. Furthermore, previous evidence indicates that excessive maternal intrauterine lipid exposures may program the development of foetus organs from early life, resulting in metabolic dysfunction. 17, 18 If maternal dyslipidaemia is a significant contributor to birth weight and implicated in neonatal metabolic dysfunction, then interventions before and during pregnancy to mitigate dyslipidaemia might improve offspring's adverse birth and metabolic health outcomes.

We performed a comprehensive systematic review and meta-analysis to explore the association, and quantify the magnitude of effect between maternal dyslipidaemia and neonatal outcomes namely, birthweight, metabolic factors, and inflammatory parameters.

Methods

Search strategy and selection criteria

The protocol for this review was registered on PROSPERO (CRD42016048568) and the review is reported in accordance with the PRISMA¹⁹ and MOOSE²⁰ guidelines. We searched systematically Embase, MEDLINE, PubMed, Scopus, CINAHL Plus, and Cochrane library (CENTRAL) up to August 1, 2016, without language or year restrictions. An updated search was made in MEDLINE before manuscript submission until the end of July 2017. The search of bibliographic databases combined index and free text terms relating to lipids (e.g. "lipids", "lipoproteins", "fatty acids", "triglycerides", "cholesterol") with those relating to pregnancy (e.g. "pregnan*", "gestation*", "gravidity", "mothers") and birthweight (e.g. "birth weight", "small for gestational age", "large for gestational age", "macrosomia"). The full strategies are provided in S1 Appendix. Cohort and Randomised Controlled Trial (RCT) filters were used to target longitudinal observational studies and the secondary analysis of RCT studies.²¹ Additional searches were conducted in Grey Literature Report and Open Grey. Reference lists of included studies were screened and checked for relevance. Search results, after removal of duplicates, were screened for relevance using title and abstract information. Fully texts of relevant articles were assessed for eligibility against the selection criteria. Screening and selection were undertaken by two reviewers independently in consultation with a third reviewer when required. This review included studies of healthy pregnant women and pregnant women with GDM or

obesity, which investigated the association between maternal lipid levels during pregnancy

authors.

(total cholesterol (TC), high-density lipoprotein cholesterol (HDL-C), low-density lipoprotein
 cholesterol (LDL-C), very-low-density lipoprotein cholesterol (VLDL-C), triglycerides (TG),
 and total free fatty acids (FFAs)) and neonatal anthropometric, metabolic, and inflammatory
 parameters.
 Studies of pregnant women with conditions that could influence maternal metabolic status

before pregnancy (hepatitis, polycystic ovary syndrome, familial hyperlipidaemia, acquired immunodeficiency syndrome, type I & type 2 diabetes, hypertension, thrombophilia, history of thromboembolism, rheumatologic disorders, cardiac dysfunction, or history of taking relevant lipid-lowering medications) were excluded.

The primary outcome was birthweight measured within the first week after delivery. Neonatal anthropometric parameters, including LBW, SGA, LGA, and macrosomia, were considered as different indexes of birthweight. Secondary outcomes included: anthropometric parameters in children less than three years old (e.g. weight gain after delivery, Body Mass Index (BMI) and skinfold thickness); biological indicators (glucose, TC, HDL-C, LDL-C, VLDL-C, TG, FFAs and insulin levels; and insulin resistance) and neonatal inflammatory factors (Monocyte Chemoattractant Protein-1 (MCP-1), interleukin 6 (IL-6), Tumour Necrosis Factor (TNF-α) and 11-beta-Hydroxysteroid Dehydrogenase Type 1 (11β HSD1) and C-reactive protein(CRP), as well as leptin levels) measured in cord blood or blood samples taken from neonates(<3 years old). Due to the diverse definition of GDM, obesity, SGA, LGA, and macrosomia in different populations, we accepted the definition specified by

Data extraction and quality assessment

A STROBE-based pre-designed form²² was used for data extraction, including the following information: study characteristics(study name, design, language, and location),

- participants(setting, eligibility/exclude criteria, and sample size), maternal characteristics (age, parity, pre-pregnancy BMI, and gestational length), follow-up (enrolment time, length of follow-up, data collection methods, and loss to follow-up rate), exposures (definition, fasting status, measured gestational weeks, and measurement methods) and outcomes (definition and measurement time point)(S2 Appendix).
- The Newcastle-Ottawa Scale was used to characterise and stratify the methodological quality of included studies (S3 Appendix). Studies quality was classified as 'low' (≤5), 'medium' (6 & 7), or 'high' (8 & 9) quality. In addition, domains relating to sample selection, comparability between groups, and method of outcome assessment were considered separately.
- Data extraction and quality assessment were conducted by two reviewers independently in consultation with a third reviewer when required (S4 Appendix). Missing information was requested from authors by email (S5 Appendix).

Data synthesis

Included studies were categorised by trimester based on the mean/median gestational age for the lipid measurement (first trimester (T1): 1-13, second trimester (T2): 14-27, and third trimester (T3): ≥28 gestation weeks). For studies reporting lipid levels multiple times within one trimester, data from the trimester with the largest sample size was adopted. Studies with different types of population (example GDM or obesity) were divided into two or three subsets to enable us to assess and report separately. Lipid measurements reported in milligrams per decilitre (mg/dL) were converted to millimoles per litre (mmol/L) using a standard unit conversion factors. ²⁴

Results of birthweight were reported in various ways, for instance, regression coefficients and correlation coefficients. Findings were summarised in tables and visually represented as horizontal histogram, displaying the direction as well as statistical significance of results comprehensively (post analysis).

Summary estimates were pooled using random effects meta-analysis, according to assessment (birthweight, SGA. LGA, and macrosomia), timing measurement(T1/T2/T3) and statistic reported in the primary study (regression coefficients, odds ratio (OR), or mean difference (MD)). Unadjusted and adjusted estimates reported in the articles were entered into random-effects models separately. Confounding factors that were adjusted (maternal age, pre-pregnancy BMI, gestational weight gain, gestational glucose level, pre-term birth, gestational lipid levels, gestational age, and neonatal gender) for each result were recorded for further sensitivity analyses. The I² statistic was used to quantify the degree of heterogeneity beyond that expected by chance in each analysis.²⁵ The potential for publication bias could not be assessed via funnel plots as the requirement for ten or more studies per meta-analysis was not met.²⁶ Due to the heterogeneity in baseline characteristics of included studies, we were not able to compare non-GDM women to GDM women. Sensitivity analysis was performed by choice of co-variates controlled for in the model. All analyses were conducted using Review Manager version 5.3 (Nordic Cochrane Centre, Copenhagen, Denmark) and R 3.3.2(The R Foundation for Statistical Computing).

Results

Study selection

Of the 13,705 unique records identified by the searches, 46 publications^{14-16, 27-69} reporting from 42 studies were included in the review (Figure 1). These studies included 31,402 pregnancies. Of the 46 included publications, 16 contributed to the quantitative analysis due

to the diversity of reporting formats (regression coefficients, correlation coefficients, mea
differences, trend analyses, or without exact effect estimates) and lack of data required for
calculations. No additional eligible studies were found in the updated search till July 2017.

Characteristics of included studies

Table describes the baseline characteristics of the 46 included publications. Most articles were published in English language and as full text articles with only one⁴⁴ study written in German, and one⁴³ published as an abstract. The studies were published between 1985 and 2016. The number of pregnancies ranged from 38 to 5,535. Based on the World Bank Income Classification of countries ⁷⁰, 25 out of 42 studies were from high income economies^{14, 16, 27, 28, 35, 41, 44-49, 51, 53, 55-62, 66-68}, 16 from upper middle economies^{15, 30-34, 37, 38, 40, 42, 43, 50, 54, 63, 65, 69}, and one middle income³⁹. Forty studies were prospective cohorts^{14, 15, 28-36, 38-50, 52, 54-57, 59-69}, three were retrospective cohorts^{27, 37, 58}, and three were secondary analyses of cohorts in RCTs^{16, 51, 53}

Quality of included studies

Forty-five publications (excluding the abstract⁴³) were assessed for methodological quality.

Ten, 21, and 14 studies were assessed as methodologically high^{15, 29, 41, 47-49, 52, 54, 60, 67},

moderate^{14, 16, 27, 30-32, 37, 38, 40, 44, 46, 50, 51, 55-57, 61, 62, 64, 68, 69} and low quality^{28, 33-36, 39, 42, 45, 53, 58, 59, 63, 65, 66} respectively(S6 Appendix). Three (7%) of 45 included studies had low risk for study selection while 40(93%) had medium risk. For comparability bias, 15(33%) had low risk, 13(29%) had medium risk, and 17(38%) had high risk. Sixteen (36%) studies were regarded to have a low risk of outcome assessment bias, with the rest (29 studies) having medium risk.

Maternal lipid levels during pregnancy and birth weight

Figure 2 shows the relationship between maternal lipid levels during pregnancy and birthweight (S7 Appendix). There were strong associations noted for HDL-C and TG

throughout pregnancy with birthweight. For HDL-C, both studies⁵⁵ reporting in T1, six^{15, 16, 31}, ^{37, 49, 55} out of 11^{15, 16, 31, 34, 37, 49, 50, 55, 59, 62} studies reporting in T2, and 11^{14, 15, 28, 41, 49, 54, 55, 61, 65}, ⁶⁸ out of 18^{14-16, 28, 39, 41, 42, 46, 49, 50, 54, 55, 58, 61, 63, 65, 68} studies reporting in T3 showed an inverse association with birthweight, while one 15 in T2 and one 16 in T3 reported a positive association. For TG, four^{52, 55, 57} out of five^{35, 52, 55, 57} studies reporting in T1, ten^{15, 31, 34, 37, 49, 55}, ^{59, 62, 67} out of 12^{15, 16, 31, 34, 37, 49, 50, 55, 59, 62, 67} studies reporting in T2, and 20^{15, 16, 39, 41, 46, 49, 50} 54-56, 58, 61, 63-65, 67, 69 out of 27^{14-16, 28, 36, 39, 41, 42, 46, 49-51, 53-56, 58, 61, 63-65, 67, 69} studies reporting in T3 found a positive association with birthweight, while three 14, 28, 51 studies in T3 reported an inverse association. Of the seven studies reporting the association between maternal FFAs level in T3 and birthweight^{36, 46, 49, 53, 56, 61, 68}, four reported a positive association^{49, 53, 56, 68}, while none reported inverse association. For TC, seven^{15, 16, 27, 37, 48, 49, 55} out of 12^{15, 16, 27, 31, 48}-50, 55, 59, 62 studies in T2, and eight^{15, 16, 48, 54-56, 65, 69} out of 22^{14-16, 28, 36, 39, 41, 42, 45, 46, 48-50, 53-56, 58,} ^{61, 63, 65, 69} studies in T3 reported a positive association, while one⁵⁵ in T2 and three^{28, 41, 55} in T3 found an inverse association. There was no evident association between maternal LDL-C level and birthweight 14, 16, 28, 31, 37, 39, 41, 42, 46, 50, 54, 55, 58, 59, 62, 63, 65, 68 or between maternal VLDL-C level and birthweight^{46, 68}. Figure 3 shows the pooled estimates for the effect of maternal lipids throughout pregnancy on birthweight using all available data (S7 Appendix). In general, the results of meta-analyses are consistent with the overall results summary (Figure 2). Maternal HDL-C was inversely associated with birthweight, particularly in T3 (adjusted RC, -70.17g per mmol/L, p<0.001). Increased maternal TG levels were significantly associated with birthweight for T1 (adjusted RC, 86.72g per mmol/L, p<0.001) and T3 (adjusted RC, 89.58g per mmol/L, p=0.01). Positive associations between TC and birthweight were observed in T1(adjusted RC, 22.67g of birthweight per mmol/L maternal lipid, p=0.02), T2 (adjusted RC, 24.74g per mmol/L, p=0.01), and T3(adjusted RC, 9.14g per mmol/L, p=0.13).

Stronger associations were observed among pregnant women with pre-pregnancy overweight or obesity in the two relevant studies (S5 Appendix).^{50, 55} The degree of heterogeneity within all meta-analyses in T3 was detected with I² values ranging from 0 to 93%. The heterogeneity decreased markedly when studies controlled for pre-pregnancy BMI, gestational weight gain, glucose level, and gestational age (S7 Appendix).

Maternal lipid levels during pregnancy and LGA, SGA, and macrosomia

Figure 4 shows the pooled adjusted OR for LGA as well as SGA, according to each type of maternal lipids in T3 (S8 & S9 Appendix). Pooled estimates for rising maternal HDL-C level revealed potentially decreased odds of LGA (OR, 0.77; 95% CI, 0.59 to 1.01; p=0.06), and significantly increased odds of SGA (OR, 1.96; 95% CI, 1.04 to 3.71; p=0.04). In contrast, increased maternal TG levels were associated with increased odds of LGA (OR, 1.08; 95% CI, 1.01 to 1.15; p=0.02), and decreased odds of SGA (OR, 0.66; 95% CI, 0.49 to 0.90; p=0.007). In addition, ten^{30, 38-40, 53, 54, 56, 58, 65, 69} out of 11^{14, 30, 38-40, 53, 54, 56, 58, 65, 69} studies reporting the association between maternal TG and LGA in T3 reported positive statistically significant associations. Of six studies investigating the relationship between maternal HDL-C and macrosomia^{30, 33, 34, 48, 47, 65}, four studies reported decreased risk of macrosomia (three statistically significant)^{30, 33, 34, 47}, especially for T2 with higher HDL-C(S10 Appendix). For the relationship of TG with macrosomia, five^{33, 38, 43, 47, 64} out of six^{30, 33, 38, 43, 47, 64} studies reported statistically significant positive OR values across three trimesters. No association was observed between maternal TC as well as LDL-C levels and LGA, SGA, and macrosomia.

Maternal lipid levels during pregnancy and other outcomes of interest

For secondary outcomes, positive correlations were found by all six publications investigating the association between different maternal lipids and different cord blood lipids, but results are inconsistent with each other ^{36, 44-46, 53, 66}. No association was observed between

maternal lipids and infant postnatal weight, weight gain, or sum of skinfolds thickness up to 2
years old^{16, 29, 51, 52}. No study investigated the relationship of maternal lipid levels during
pregnancy with neonatal glucose, insulin, inflammatory factors and leptin levels in our
searches.

Discussion

Summary of the findings

This is the first systematic review pooling data from 40 longitudinal observational studies and two RCT secondary analysis studies providing quantitative estimates of the magnitude of association between maternal lipid levels at various stages of pregnancy and neonatal health outcomes. Throughout pregnancy, low maternal HDL-C and high TG levels are associated with increased birthweight. Low HDL-C and high TG increased the risk of LGA/macrosomia and lowered the risk of SGA babies. Maternal TC level throughout pregnancy and FFAs level in the third trimester are positively associated with a small increase in birthweight. Associations are stronger among populations with pre-pregnancy obesity. The findings provide evidence for the critical role of dyslipidaemia in gestational metabolism and neonatal health, and will contribute to future research and management of gestational dyslipidaemia.

Potential mechanisms

The results are mostly consistent with previous published evidence. Maternal lipid metabolism is mainly in lipogenesis state in the earlier half of pregnancy, but then switches into catabolic state.^{71,72} When the lipid accumulation exceeds the storage capacity of adipose tissue, the buffering function of the adipocytes is decreased, leading to elevated serum FFAs and TG.⁷³⁻⁷⁵ Compared to pregnant women with smaller pre-pregnancy BMI, women who are overweight or obese will not only progress to catabolic state earlier, but also have less capacity to inhibit lipolysis.¹⁸ Women with obesity prior to pregnancy usually present with

292	more central adipose accumulation and severe dyslipidaemia ^{76, 77} , resulting in steep
293	concentration gradient across the placenta. ⁷⁸
294	Both in vivo and epidemiological evidence suggest that excessive maternal intrauterine lipid
295	exposure could affect the development of foetus organs systematically, which can then alter
296	initial foetus metabolism and feeding behaviours permanently. ^{18, 79} Previous animal studies
297	observed that foetal metabolic abnormalities mediated by maternal obesity and high-fat diet
298	often manifest as increased body weight, fat mass, blood glucose, cholesterol and blood
299	pressure levels; and decreased insulin sensitivity and ectopic lipid storage in newborns. 18 The
300	latest multi-ancestry genome-wide association study (GWAS) meta-analysis also
301	demonstrated that cholesterol biosynthesis is one of the most important metabolic pathways
302	involved in birthweight. 17 Strengths and weakness
303	The major strengths of this study are the comprehensive searches, adherence to robust review
304	methodology and thorough analyses. Special care was taken in the handling of missing data,
305	which was addressed by personal contact with the authors in an attempt to minimise reporting
306	bias. The inclusion of longitudinal studies ensured the temporal association between
307	exposures and outcomes, which also permitted a trimester-specific analysis. The major
308	limitation of the study was the substantial heterogeneity, possibly due to the diversity of
309	settings, study populations, lipid measurement methods and diverse gestational age of the
310	studied populations. However, this heterogeneity was addressed by subgroup analysis.
311	It would be intriguing to explore the effects of maternal dyslipidaemia independent of
312	maternal hyperglycaemia. Unfortunately, this was not feasible due to the nature of data
313	reported in individual study. GDM women are known to have higher TG levels and lower
314	HDL-C levels compared with non-GDM women. ¹¹ However, elevated maternal TG levels
315	and lower HDL-C levels are associated with the risk of LGA and macrosomia in both GDM

women^{38, 53, 58} and non-GDM women^{30, 39, 40, 52, 54, 69}. For women with type 1 diabetes/GDM, maternal hyperglycaemia is not the sole contributor to increased birth weight since foetuses may develop LGA despite them having optimal glycaemic control. 80 Several other studies found that lipid levels during pregnancy, similar to glucose levels, are also strong metabolic determinants for foetal growth 15, 29, 31, 32, 35, 37, 41, 47, 53, 56, 61, 64. Our sensitivity analyses result also shown there is little effect on the relationship between gestational HDL-C/TG levels and birth weight when removing those studies controlled for glucose (S7.13 & S7.23). Collectively, this evidence suggests that maternal dyslipidaemia may be an independent, unrecognised risk factor of LGA/macrosomia. Unfortunately, paucity of the required primary data prevented the pre-specified subgroup analyses on the basis of different definitions used for GDM and obesity across studies. Thus, this should be acknowledged as a source of clinical heterogeneity when interpreting the findings of the present study. Another limitation of this study is that we are unable to control for the effect of GDM treatment on lipid levels. However, it has been noticed that initiation of therapy (diet control, insulin, or metformin) may modestly influence TG levels⁸¹, yet to a direction that would obscure rather than magnify differences between normal and GDM pregnancies. Similarly, our sensitivity analyses shown a moderate decrease on triglycerides effect estimate when removing studies that excluded pre-term births (\$7.25). It should be acknowledged that our primary outcome, birth weight, is a quite inexact measure of foetal growth, although it has been widely measured and utilized in clinical and research areas. We tried to extend our target outcomes from birth weight parameters to other neonatal growth parameters, biological indicators, and inflammatory factors, however, we did not find sufficient studies.

Implications

Our results provides compelling evidence on the role of maternal circulating HDL-C and TG levels on birth outcomes, and suggest that the under-recognised adverse effects of intrauterine exposure to maternal dyslipidaemia may need further investigation in large prospective cohorts or in randomised trials. Although the importance of screening for preconceptional dyslipidaemia has been noted in recent guidelines to alert for risk assessment for GDM^{82, 83}, its independent adverse effects remain largely underestimated in routine clinical practice and recommendations regarding the management of dyslipidaemia preconceptionally or during pregnancy are still lacking. Our findings do question the current clinical practice and support the monitoring of gestational dyslipidaemia before or during pregnancy. Moreover, our findings may be a call for action regarding the implementation of strategies to address maternal dyslipidaemia (such as carefully planned dietary interventions, increasing physical activity, and/or Omega-3 fatty acids supplementation). Meanwhile, gestational dyslipidaemia, as an important feature of obesity and GDM, might be a potential treatment target for clinical interventions. These steps need to be evaluated by global health policy makers through randomised controlled trials, evidence synthesis and consensus. ⁸⁴⁻⁸⁶

Conclusion

Our findings demonstrate that maternal low HDL-C and high TG levels are positively associated with neonatal birthweight. No effect was documented for total or LDL cholesterol. Findings are of clinical importance in considering the management of gestational dyslipidaemia, for example using lifestyle interventions and omega-3 fatty acid supplementation to improve maternal and neonatal outcomes.

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Contributors: KN, QX, KK, and JW conceived the research question. JW defined the question, designed the study, and conducted searches, data extraction, quality assessment, and data analysis. AS and KN contributed as second reviewers for the data extraction and quality assessment. DM, KN, and MJP advised on study design and contributed to data analysis. KK, QX, PS, and KAT also provided input for study design. All authors contributed to the interpretation of the results. JW led the writing of the manuscript with critical input from all other authors. All authors, external and internal, had full access to all data (including statistical reports and tables) in the study and can take responsibility for the integrity of the data and the accuracy of the data analysis. JW is the guarantor.

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Competing interests: All authors have completed the ICMJE uniform disclosure form at www.icmje.org/coi_disclosure.pdf and declare: no support from any organisation for the submitted work; no financial relationships with any organisations that might have an interest in the submitted work in the previous three years; no other relationships or activities that could appear to have influenced the submitted work.

- Ethical Approval: Not required.
- **Data sharing:** No additional data available.

Transparency: The lead author (JW) affirms that this manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as planned (and, if relevant, registered) have been explained.



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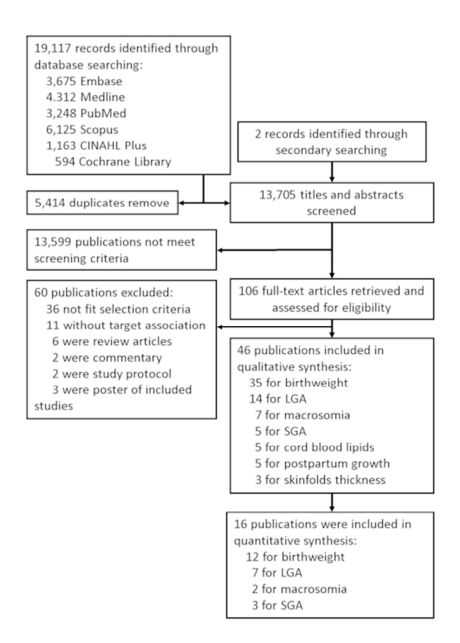
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625 Table Baseline characteristics of included studies

Study ID	Study design	Locations	Population (N)	TC	HDL	LDL	TG	VLDL	FFAs	Tri.	Outcomes
Ye et al.2015 ⁵⁴	Prospective observational study	China	non-GDM (n=1,243)	1	V	V	V			3	Birthweight LGA, SGA
Wang et al.2015 ³¹	Prospective cohort study	China	General (n=636)		\checkmark	$\sqrt{}$				2	Birthweight
Crume et al.2015 ⁴⁹	Prospective cohort study	American	General (n=804)		\checkmark				$\sqrt{}$	2,3	Birthweight
Hwang et al.2015 ⁶⁷	Prospective cohort study	Korea	non-GDM (n=1,011)				$\sqrt{}$			2,3	Birthweight
Kulkarni et al.2013 ¹⁵	Prospective cohort study	India	non-GDM (n=631)		$\sqrt{}$		$\sqrt{}$			2,3	Birthweight
Vrijkotte et al.2012 ⁵²	Prospective cohort study	Netherlands	non-GDM (n=4,008)				$\sqrt{}$			1	LGA, SGA
Retnakaran et al.2012 ¹⁴	study	Canada	non-GDM (n=472)		$\sqrt{}$	$\sqrt{}$				3	Birthweight LGA
Hou et al.2014 ⁴⁰	Prospective observational study	China	non-GDM (n=2,790)	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$			3	LGA
Kramer et al.2014 ²⁹	Prospective cohort study	Canada	General (n=340)		$\sqrt{}$		$\sqrt{}$			3	Infant weight gain at 3 months
Son et al.2010 ⁵⁸	Retrospective longitudinal observational study	Korea	GDM (n=104)	V	$\sqrt{}$	V	$\sqrt{}$			3	Birthweight LGA
Ahmad et al. 2006 ⁶⁹	Controlled prospective study	Malaysia	non-GDM (n=246)							3	Birthweight LGA
Di et al. 2005 ⁵⁹	Prospective observational study	Italy	OGTT+ (n=83)		\checkmark	$\sqrt{}$	$\sqrt{}$			2	Birthweight LGA
Couch et al.1998(1) ⁴⁶ Couch et al.1998(2) ⁴⁶	Prospective observational study	American	GDM (n=20) Non-GDM (n=20)	1	V	√	V	V	$\sqrt{}$	3	Birthweight Cord vein lipids profile
Ortega et al. 1996 ⁴⁵	Prospective cohort study	Spain	General (n=292)	1	\checkmark	$\sqrt{}$	1	$\sqrt{}$	V	3	Birthweight Cord arteriovenous lipids profile
Alberti-Fidanza et al. 1995 ⁶⁶	Prospective observational study	Italy	General (n=70)	1	V					1-3	Mixed venous- arterial cord blood lipids profile
Schaefer-Graf et al. 2008 ⁵³	Secondary analysis of RCT study	German	GDM (n=150)	V			$\sqrt{}$		V	3	Birthweight, cord blood lipids LGA
Swierzewska et al. 2015 ⁴²	Prospective observational study	Poland	General (n=136)		$\sqrt{}$	V				3	Birthweight
Sommer et al. 2015 ⁴¹	Prospective cohort study	Norway	General (n=699)		$\sqrt{}$	$\sqrt{}$	$\sqrt{}$			3	Birthweight, sum of skinfolds
Slagjana et al. 2014 ³⁹	Prospective cohort study	Yugoslavia	non-GDM (n=200)		\checkmark	$\sqrt{}$	$\sqrt{}$			3	Birthweight LGA, SGA
Laleh et al. 2013 ³⁸	Prospective cohort study	Iran	GDM (n=112)		$\sqrt{}$	V	$\sqrt{}$			3	LGA, macrosomia
Whyte et al. 2013 ⁶²	Prospective cohort study	Ireland	General (n=189)		\checkmark	$\sqrt{}$				2	Birthweight
Zhou et al. 2012 ³³	Prospective cohort study	China	General (n=1,000)	1	$\sqrt{}$	V	$\sqrt{}$			2	Macrosomia
Vrijkotte et al. 2011 ⁶⁰	Prospective cohort study	Netherlands	General (n=2,052)	V			V			1	Birthweight Postpartum growth
Vinod et al.2011(1) ⁵⁵ Vinod et al.2011(2) ⁵⁵	Prospective cohort study	American	Overweight (n=71) Normal weight (n=72)	√	$\sqrt{}$	√	V			1-3	Birthweight
Zawiejska et al.	Prospective	Poland	GDM		√		$\sqrt{}$			2	Birthweight

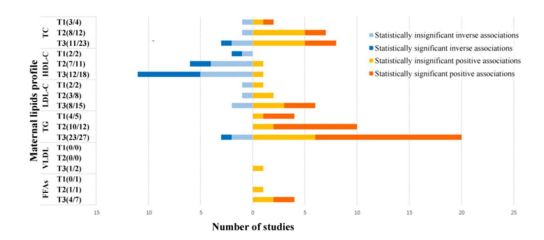
Study ID	Study design	Locations	Population (N)	TC	HDL	LDL	TG	VLDL	FFAs	Tri.	Outcomes
2008 ³⁴	observational study		(n=357)								Macrosomia
Clausen et al. 2005 ⁴⁷	Prospective cohort study	Norway	General (n=2,050)		$\sqrt{}$	$\sqrt{}$				2	Macrosomia
Mathews et al. 2003 ⁴⁸	Prospective cohort study	UK	General (n=798)							2,3	Birthweight
Olmos et al.2014(1) ⁵⁰ Olmos et al.2014(2) ⁵⁰ Olmos et al.2014(3) ⁵⁰	Prospective observational study	Chile	GDM + lean (n=128) GDM + overweight (n=105) GDM + obese (n=46)	√	1		V			2,3	Birthweight
Emet et al.2013 ⁶³	Prospective observational study	Turkey	General (n=801)		$\sqrt{}$	$\sqrt{}$				3	Birthweight, infant weight at 3 months
Liu et al.2016 ³⁷	Retrospective cohort study	China	General (n=1,546)		$\sqrt{}$	$\sqrt{}$				2	Birthweight
Brunner et al. 2013 ⁵¹	Secondary analyses of RCT study	German	General (n=208)				V			3	Birthweight, postpartum growth, skinfolds thickness
Knopp et al.1992 ⁶⁴	Prospective observational study	American	NS- (n=521) PS+ (n=264) GDM (n=96)				V			3	Birthweight
Knopp et al.1985 ⁶⁸	Prospective observational study	American	General (n=283)		$\sqrt{}$	$\sqrt{}$		\checkmark	$\sqrt{}$	3	Birthweight
Schaefer-Graf et al. 2011 ³⁶	Prospective observational study	German	non-GDM (n=190)	V			V		$\sqrt{}$	3	Birthweight, Cord blood metabolic parameters
Nolan et al.1995 ⁵⁷	Prospective observational study	Australia	General (n=388)							1	Birthweight
Lin et al.2013 ⁴³	Prospective observational study	China	General (ND)							ND	Macrosomia
Friis et al.2012 ⁶¹	Prospective observational study	Norway	General (n=207)	V	√				$\sqrt{}$	3	Birthweight
Lei et al.2016 ³²	Prospective cohort study	China	General (n=5,535)		$\sqrt{}$					2	LGA, SGA
Kitajima et al. 2001 ⁵⁶	Prospective observational study	Japan	OGTT + (n=146)	1			1		$\sqrt{}$	3	Birthweight LGA
Mossayebi et al. 2014 ⁶⁵	Prospective cohort study	Iran	General (n=154)	√	√	$\sqrt{}$	1			3	Birthweight LGA, macrosomia
Geraghty et al. 2016 ¹⁶	RC1 study	UK	non-GDM (n=331)		$\sqrt{}$	$\sqrt{}$	V			2,3	Birthweight Postpartum growth, sum of skinfolds
Jin et al. 2016 ³⁰	Prospective cohort study	China	non-GDM (n=934)			$\sqrt{}$				1-3	LGA, SGA, macrosomia
Brockerhoff 1986 ⁴⁴	Prospective observational study	German	ND (n=112)		$\sqrt{}$	$\sqrt{}$		$\sqrt{}$		2	Cord blood lipids profile
Harmon et al. 2011 ³⁵	Prospective observational study	American	non-GDM (n=38)						$\sqrt{}$	1	Birthweight
Robin et al. 2007 ²⁷	Retrospective cohort study	American	General (n=957)							2	Birthweight
Charles et al. 2016 ²⁸	Perspective observational study	Mediterranean countries	General (n=1062)	\checkmark	$\sqrt{}$	$\sqrt{}$				3	Birthweight

Abbreviation: Trimester(Tri), Total cholesterol (TC), high-density lipoprotein cholesterol (HDL-C), low-density lipoprotein cholesterol (LDL-C), very low-density lipoprotein cholesterol (VLDL-C), triglycerides(TG), free fatty acids(FFAs), large-for-gestational age(LGA), small for gestational age(SGA), randomized controlled trial(RCT), and no documented(ND).



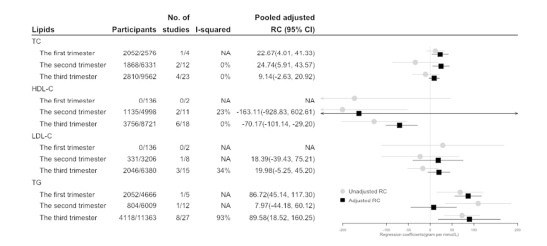
Title: Figure 1. Flow-diagram of study selection

171x244mm (72 x 72 DPI)



Title: Figure 2. Results summary of the association of maternal lipid levels with birth weight throughout pregnancy!! + Notes: The numbers in parenthesis: The number of studies shown in this figure/the overall number of studies reporting the target associations. Studies reporting statistically insignificant results without its direction or those that did not report their results are not shown in the figure.

253x115mm (72 x 72 DPI)

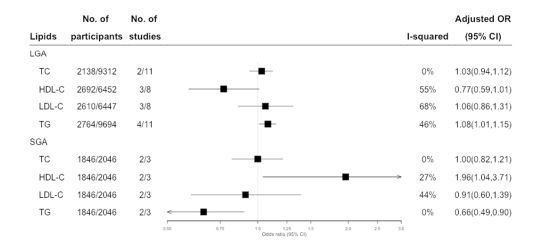


Title: Figure 3 Summary of findings of meta-analysis for the associations between maternal lipids and birth weight throughout pregnancy

Notes: The number of participants (studies) included into quantitative analysis/ overall number of participants (studies) that reported the outcome of interest.

The number of participants (studies) included into quantitative analysis/ overall number of participants (studies) that reported the outcome of interest.

339x166mm (72 x 72 DPI)



Title: Figure 4 Summary of findings of meta-analysis for the associations between maternal lipids and LGA/ SGA in the third trimester

Notes: The number of participants (studies) included into quantitative analysis/ overall number of participants (studies) that reported the outcome of interest.

336x166mm (72 x 72 DPI)

Policy.

Supplementary material

Context

S1 Appendix Sample search in Medline	4
S2 Appendix Data extraction form	5
S3 Appendix Newcastle-Ottawa Scale	7
S4 Appendix Basic characteristics extraction form	
S5 Appendix Results extraction form	
S6 Appendix Quality assessment form	
S7 Appendix Data analysis for birthweight	
Data summary	
S7.1 Table Results summary of the association of maternal lipid levels with birthweight throughout pregna	
Total cholesterol (TC)	•
S7.2 Table Results summary of the association of maternal TC level with birthweight	
Meta-analysis	
S7.1 Figure Overall meta-analysis of crude regression coefficients for the association between materna levels and birthweight throughout pregnancy	
S7.2 Figure Overall meta-analysis of adjusted regression coefficients for the association between mater levels and birthweight throughout pregnancy	
Subgroup analysis	55
S7.3 Figure Adjusted regression coefficient_General vs. non-GDM_the 2nd trimester_Random effect to	
S7.4 Figure Adjusted regression coefficient_ General vs. non-GDM_ the 3 rd trimester_ Random effect m	
Sensitivity analysis	
S7.5 Figure Adjusted regression coefficients_ exclude studies control for pre-pregnancy BMI or gestation weight gain	
S7.6 Figure Adjusted regression coefficients_ exclude studies control for maternal glucose level	56
S7.7 Figure Crude regression coefficients_ exclude studies control for pre-term birth	57
S7.8 Figure Adjusted regression coefficients_ exclude studies that did not control for pre-term birth	
High-Density lipoprotein Cholesterol (HDL-C)	
S7.3 Table Results summary of the association of maternal HDL-C level with birthweight	
Meta-analysis	59
S7.9 Figure Overall meta-analysis of crude regression coefficients for the association between materna levels and birthweight throughout pregnancy	
S7.10 Figure Overall meta-analysis of adjusted regression coefficients for the association between mate HDL-C levels and birthweight throughout pregnancy	
Subgroup analysis	
S7.11 Figure Adjusted regression coefficient_ General vs. non-GDM_ the 3rd trimester_ Random effect	
Sensitivity analysis	60
S7.12 Figure Adjusted regression coefficients_ exclude studies control for pre-pregnancy BMI or gestat weight gain	
S7.13 Figure Adjusted regression coefficients_ exclude studies control for maternal glucose level	60
S7.14 Figure Adjusted regression coefficients_ exclude studies control for pre-term birth	61
Low-Density lipoprotein Cholesterol (LDL-C)	62
S7.4 Table Results summary of the association of maternal LDL-C level with birthweight	
Meta-analysis	
S7.15 Figure Overall meta-analysis of crude regression coefficients for the association between matern C levels and birthweight throughout pregnancy	63
S7.16 Figure Overall meta-analysis of adjusted regression coefficients for the association between mate LDL-C levels and birthweight throughout pregnancy	

Sensitivity analysis	64
S7.17 Figure Adjusted regression coefficients_ exclude studies that did not control for pre-term	n birth64
S7.18 Figure Adjusted regression coefficients_ exclude studies that did not control for other m levels	•
Triglycerides (TG)	65
S7.5 Table Results summary of the association of maternal TG level with birthweight	65
Meta-analysis	67
S7.19 Figure Overall meta-analysis of crude regression coefficients for the association betwee levels and birthweight throughout pregnancy	
S7.20 Figure Overall meta-analysis of adjusted regression coefficients for the association between levels and birthweight throughout pregnancy	
Subgroup analysis	68
S7.21 Figure Adjusted regression coefficient_ General vs. non-GDM_ the 3rd trimester_ Rand	***
Sensitivity analysis	
S7.22 Figure Adjusted regression coefficients_ the 3rd trimester_ exclude studies control for p BMI or gestational weight gain	
S7.23 Figure Adjusted regression coefficients_ the 3rd trimester_ exclude studies control for n level	0
S7.24 Figure Adjusted regression coefficients_ the 3rd trimester_ exclude studies control for o lipid levels	
S7.25 Figure Adjusted regression coefficients_ the 3rd trimester_ exclude studies control for p S7.26 Figure Adjusted regression coefficients_ the 3rd trimester_ exclude studies that did not gestational age	control for
Free Fatty Acids (FFAs)	70
S7.6 Table Results summary of the association of maternal FFAs levels with birthweight	70
Very Low-density lipoprotein cholesterol (VLDL)	70
S7.7 Table Results summary of the association of maternal VLDL-C levels with birthweight	70
upplementary 8 Data analysis for Large for gestational age	71
Total cholesterol (TC)	
S8.1 Table Results summary of the association of maternal TC levels with LGA	
Meta-analysis.	
S8.1 Figure Meta-analysis of adjusted odds ratio for the association between maternal TC leve	
S8.2 Figure Meta-analysis for mean difference of maternal TC levels between LGA and referent third trimester	nce groups in the
High-density lipoprotein cholesterol (HDL-C)	
S8.2 Table Results summary of the association of maternal HDL-C levels with LGA	
Meta-analysis	
S8.3 Figure Meta-analysis of adjusted odds ratio for the association between maternal HDL-C in the third trimester	C levels and LGA
S8.4 Figure Meta-analysis for mean difference of maternal HDL-C levels between LGA and re the third trimester	eference groups in
Sensitivity analysis	74
S8.5 Figure Sensitivity analysis_ Adjusted odds ratio_ Exclude study adjust for other maternal	l lipid levels 74
Low-density lipoprotein cholesterol (LDL-C)	75
S8.3 Table Results summary of the association of maternal LDL-C levels with LGA	
Meta-analysis	76
S8.4 Figure Meta-analysis of adjusted odds ratio for the association between maternal LDL-C in the third trimester	<i>76</i>

Triglycerides (TG)	77
S8.4 Table Results summary of the association of maternal TG levels with LGA	77
Meta-analysis	78
S8.6 Figure Meta-analysis of adjusted odds ratio for the association between maternal TG levels and LGA throughout pregnancy	
S8.7 Figure Forest plots of crude odds ratio for the association between maternal TG levels and LGA throughout pregnancy	78
S8.8 Figure Forest plots of adjusted odds ratio for the association between maternal TG levels and LGA throughout pregnancy	79
Sensitivity ananlysis	79
S8.9 Figure Sensitivity analysis_ Exclude studies adjust for other maternal lipid levels	79
Free fatty acids (FFAs)	80
S8.5 Table Results summary of the association of maternal FFAs levels with LGA	80
Supplementary 9 Data analysis for Small for gestational age (SGA)	81
Total cholesterol (TC)	81
S9.1 Table Results summary of the association of maternal TC levels with SGA	81
S9.1 Figure Meta-analysis of adjusted odds ratio for the association between maternal TC levels and SGA throughout pregnancy	81
High-density lipoprotein cholesterol (HDL-C)	82
S9.2 Table Results summary of the association of maternal HDL-C levels with SGA	82
S9.2 Figure Meta-analysis of adjusted odds ratio for the association between maternal HDL-C levels and SGA throughout pregnancy	
Low-density lipoprotein cholesterol (LDL-C)	83
S9.3 Table Results summary of the association of maternal LDL-C levels with SGA	83
S9.3 Figure Meta-analysis of adjusted odds ratio for the association between maternal LDL-C levels and SGA the third trimester	
Triglycerides (TG)	84
S9.4 Table Results summary of the association of maternal TG levels with SGA	84
S9.4 Figure Meta-analysis of adjusted odds ratio for the association between maternal TG levels and SGA throughout pregnancy	84
Supplementary 10 Data analysis for Macrosomia	85
Total cholesterol (TC)	
S10.1 Table Results summary of the association of maternal TC levels with macrosomia	
High-density lipoprotein cholesterol (HDL-C)	
S10.2 Table Results summary of the association of maternal HDL-C levels with macrosomia	
S10.1 Figure Forest plots of adjusted odds ratio for the association between maternal HDL-C levels and macrosomia throughout pregnancy	
Low-density lipoprotein cholesterol (LDL-C)	
S10.3 Table Results summary of the association of maternal LDL-C levels with macrosomia	
Triglycerides (TG)	
S10.4 Table Results summary of the association of maternal TG levels with macrosomia	
S10.2 Figure Meta-analysis of adjusted odds ratio for the association between maternal TG levels and macros	omia
\$10.3 Figure Forest plots of adjusted odds ratio for the association between maternal TG levels and macrosor	

1 2 S1 Appendix Sample search in Medline 3 1. exp Lipids/ or lipid\$.mp. 4 2. lipoprotein\$.mp. or exp Lipoproteins/ 5 3. exp Fatty Acids/ or fat* acids.mp. 4. triglycerides.mp. or exp Triglycerides/ 6 5. exp Lipoproteins, VLDL/ or exp Cholesterol, VLDL/ or VLDL.mp. 7 6. LDL.mp. or exp Cholesterol, LDL/ or exp Lipoproteins, LDL/ 8 7. IDL.mp. or exp Lipoproteins, IDL/ 9 8. exp Lipoproteins, HDL/ or exp Cholesterol, HDL/ or HDL.mp. 10 9. exp Cholesterol/ or cholesterol.mp. or exp Cholesterol Esters/ 11 10. hyperlipid?emia\$.mp. or exp Hyperlipidemias/ 12 11. dyslipid?emia\$.mp. or exp Dyslipidemias/ 13 12. hypertriglycerid?emia\$.mp. or exp Hypertriglyceridemia/ 14 13. hypercholesterol?emia.mp. or exp Hypercholesterolemia/ 15 14. metabolic.mp. 16 15. 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12 or 13 or 14 17 16. exp Maternal Health/ or maternal.mp. 18 17. exp Pregnanes/ or pregnan*.mp. 19 18. exp Pregnancy/ or gestation*.mp. 20 19. gravidity.mp. or exp Gravidity/ 21 20. mother\$.mp. or exp Mothers/ 22 21. 16 or 17 or 18 or 19 or 20 23 22. (birth weight or birthweight).mp. or exp Birth Weight/ or exp Infant, Low Birth Weight/ 24 23. overweight.mp. or exp Obesity/ or exp Overweight/ or exp Body Weight/ 25 24. (SGA or Small for gestational age).mp. or exp Infant, Small for Gestational Age/ 26 25. (LGA or Large for gestational age).mp. 27 26. exp Fetal Macrosomia/ or macrosomia.mp. 27. exp "Growth and Development"/ or exp Growth/ or (growth or development).mp. or exp Fetal Growth Retardation/ 28 29 or exp Fetal Development/ or exp Child Development/ 28. weight gain.mp. or exp Weight Gain/ 30 29. (hyperglyc?emia or hypoglyc?emia).mp. or exp Hyperglycemia/ or exp Hypoglycemia/ 31 30. (insulin* or hyperinsulinism or IR).mp. or exp Insulin/ or exp Insulin Resistance/ or exp Hyperinsulinism/ 32 31. exp Glucose Intolerance/ or glucose.mp. or exp Glucose/ or exp Glucose Metabolism Disorders/ 33 32. skinfold thickness.mp. or exp Skinfold Thickness/ 34 33. (monocyte chemoattractant protein-1 or MCP-1).mp. 35 34. (interleukin 6 or IL-6).mp. 36 35. exp Tumor Necrosis Factor-alpha/ or tumour necrosis factor-alpha.mp. 37 36. exp 11-beta-Hydroxysteroid Dehydrogenase Type 1/ or HSD1.mp. 38 37. exp Leptin/ or leptin.mp. 39 38. exp Inflammation/ or inflammat*.mp. 40 39. 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12 or 13 or 14 or 22 or 23 or 24 or 25 or 26 or 27 or 28 or 29 or 41 30 or 31 or 32 or 33 or 34 or 35 or 36 or 37 or 38 42 40. (neonatal or fetal or foetal or fetus or foetus or infant or offspring or new born).mp. or exp Infant/ 43 41. 15 and 21 and (39 and 40) 44 42. (animal or mouse or mice or rodent or sheep or mutton or pig or hoggory or hog or swine or rabbit\$).mp. 45 43. 41 not 42 46 44. cohort studies/ or longitudinal studies/ or follow-up studies/ or prospective studies/ or retrospective studies/ or 47 cohort.ti,ab. or longitudinal.ti,ab. or prospective.ti,ab. or retrospective.ti,ab. 48

45. "randomized controlled trial".pt.

46. (random\$ or placebo\$ or single blind\$ or double blind\$ or triple blind\$).ti,ab.

47. (retraction of publication or retracted publication).pt.

48. or/44-47

49

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51 52

53

54

55

56

57

58

59 60 49. (animals not humans).sh.

50. ((comment or editorial or meta-analysis or practice-guideline or review or letter or journal correspondence) not "randomized controlled trial").pt.

51. (random sampl\$ or random digit\$ or random effect\$ or random survey or random regression).ti,ab. not "randomized controlled trial".pt.

52. or/49-51

53. 48 not 52

54. 43 and 53

S2 Appendix Data extraction form

A. Reference information

- 1. ID number
- 2. Title
- 3. Author
- 4. Journal
- 5. Publication Year
- 6. Language
- 7. Sponsor

B. Study design

- 1. Study design
- 2. Setting
- 3. Locations
- 4. Data collection

C. Participants

- 1. Eligibility criteria (source and methods of selection of participants)
- 2. Matching criteria (if applicable)
 - a. Matching criteria
- b. Attempts were made within the design or analysis to balance the comparison groups for potential confounders (YES/NO).
- c. The groups are comparable at baseline, including all major confounding and prognostic factors (YES/NO).
- 3. Sample Size
 - a. Number of both exposed and unexposed groups
 - b. Report numbers of individuals at each stage of study
 - c. Give reasons for non-participation at each stage (YES/NO)
 - d. Does the size of samples have enough power to detect the difference of primary outcomes?
- 4. Demographic, clinical and social characteristics
 - a. Age
 - b. Ethnicity
 - c. Pre-pregnant BMI/weight
 - d. Marital status
 - e. Education
 - f. Other potential confounders information

D. Follow-up

- 1. Enrolment time
- 2. Length of follow-up
 - a. Length of follow-up (average and total amount) $\,$
- b. All groups were followed up for an equal length of time (or analysis was adjusted to allow for differences in length of follow-up)
- 3. Methods of follow-up
- 4. Lost to follow-up
 - a. Attrition rate in each group
 - b. How many participants in each group were no outcome data available? (number & proportion)
 - c. Does it comparable? (YES/NO)

E. Exposure

- 1. Definition of exposures
- 2. When did they take samples
- 3. Exposure measurement

F. Outcomes

- 1. Primary outcomes (definition and measurement)
- 2. Secondary outcomes (definition and measurement)

G. Statistical methods

- 1. Statistical methods, including those used to control for confounding
- 2. Describe any methods used to examine subgroups and interactions
- 3. How missing data were addressed
- 4. Explain how lost to follow-up was addressed
- 5. Describe any sensitivity analysis

H. Results

60

1. Number of outcomes events or summary measures over time

- 2. Give unadjusted estimates and, if applicable, confound der-adjusted estimates and their precision (e.g. 95% confidence interval). Make clear which confounders were adjusted for and why they were included
 - 3. Report category boundaries when continuous variables were categorized
 - 4. Alpha value and beta value

I. Limitations

1. Interpretation

Give a cautious overall interpretation of results considering objectives, limitations, multiplicity of analyses, results from similar studies, and other relevant evidence.

2. Generalizability (external validity)

J. Other notes



S3 Appendix Newcastle-Ottawa Scale

Selection

- 1. Representativeness of exposed cohort population
 - 1) Truly representative of the average, community-dwelling target pregnant women \star
 - 2) Somewhat representative of the average, community-dwelling target pregnant women★
 - 3) Selected group of pregnant women, e.g. only certain socio-economic groups/areas
 - 4) No description of the derivation of the cohort
- 2. Selection of the unexposed cohort
 - 1) Drawn from the same source as the exposed cohort★
 - 2) Drawn from a different source
 - 3) No description of the derivation of the unexposed cohort
- 3. Ascertainment of exposures
 - 1) Laboratory diagnosed ★
 - 2) Secure record (e.g. health care/clinical record) ★
 - 3) Written self-report
 - 4) Other/ no description
- 4. Demonstration that outcome of interest was not present at start of study
 - 1) Yes★
 - 2) No

Comparability

- 1. Comparability of cohort based on the design or analysis
 - 1) Study controls for
 - ① Outcomes measured at delivery: gestational age *
 - 2 Outcomes measured over 1 month after delivery: neonatal age *
- 2) Study controls for any two of additional factors (e.g. neonatal gender, maternal age, parity, socio-economic level, cigarette exposures, delivery mode and so on)

Outcome

- 1. Assessment of outcomes
 - 1) Independent blind assessment★
 - 2) Record linkage★
 - 3) Self-report
 - 4) Other/ no description
- 2. Was follow up long enough for outcomes to occur
 - 1) Yes, if the study follow their subjects until outcomes occur★
 - 2) No, if the study follow their subjects until outcomes occur
- 3. Adequacy of follow up of cohorts
 - 1) Complete follow up: all subjects accounted for★
- 2) Subjects lost to follow up unlikely to introduce bias: number lost \leq 20%, or description of those lost suggesting no different from those followed \bigstar
 - 3) Follow up rate <80% and no description of those lost
 - 4) No statement

S4 Appendix Basic characteristics extraction form

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
Ye et al. 2015	Study design: Prospective observational study Language: English Location: China	Setting: Maternal and Child Health centres (MCH) of Hefei. Eligibility criteria: Women (≥18 years) who given birth in MCH centres of Hefei around 36 th – 41 st gestation week. Exclude criteria: 1) Gestational diabetes, overt diabetes, hypertension and heart disease. 2) Preterm births (before 37 weeks) or multiple pregnancies. 3) No information on birth weight. Sample size: n=1,243	$\overline{\mathbf{x} \pm \mathbf{SD}}$ or \mathbf{n} (%) $\frac{Age (year)}{27.9 \pm 4.3}$ $\frac{Primiparous}{1012 (81.4)}$ $\frac{Pre-pregnancy}{BMI (kg/m^2)}$ 20.5 ± 2.5 $\frac{Gestational \ length}{39.6 \pm 1.0}$ $\frac{Fasting \ blood}{No \ Statement}$	Enrolment time Gestational age at entry (36 th – 41 st gestation week) (1 st Jan 2011 – 31 st July 2012) Length Follow up until birth Methods Clinical follow-up Data collection Questionnaire, clinical medical records Loss to follow-up 0	Maternal serum TG, TC, HDL, LDL were measured close to delivery (36-41 weeks, in most case 1 week to delivery)	Birth weight was retrieved from medical records after delivery. LGA: infants with birth weight > 90 th percentile for local population after adjusting for gestational age and sex. SGA: birth weight < 10 th AGA: 10 th Sbirth weight < 90 th	8
Wang et al. 2015	Study design: Cohort Language: English Location: China	Setting: No statement Eligibility criteria: 1) Chinese women with a singleton pregnancy and a live delivery; 2) have GDM screening at 24-28 weeks of gestation; 3) presented for booking at or before 16 weeks and gave birth at or after 36 weeks; 4) compete antenatal and birth data. Exclude criteria: Type 1 or type 2 diabetes; hyperlipidaemia, hypertension, cardiovascular diseases or metabolic syndrome before pregnancy; a history of severe systemic disease (liver cirrhosis, chronic renal failure, severe anaemia or immune disorders); and untreated endocrinopathies (hyperadrenalism, hypoadrenalism, hyperthyroidism or hypothyroidism) Sample size: n= 636	Median (25 th -75 th) Age (year) Non-GDM: 29 (27-31) GDM: 31 (29-34) Parity No statement Pre-pregnancy BMI (kg/m²) Non-GDM: 20.03 (18.59- 21.55) GDM: 21.02 (19.24- 22.56) Gestational length Non-GDM: 39 (39-40) GDM: 39 (38-40) Fasting blood Yes.	Enrolment time: Gestational age at entry (at or before 16 th gestation week) (1 st Jan 2013 – 31 st Dec 2013) Length: At least follow up until birth Methods: No statement Data collection: laboratory diagnosis Loss to follow-up: 0	Maternal overnight fasting blood was taken at the time of OGTT (24 th -28 th weeks) for TC, HDL, LDL and TGs laboratory analyses (standard enzymatic procedures on automatic chemistry analyser).	Birthweight.	6

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		(110 GDM and 526 non-GDM)					
Crume et al.2015	Study design: Prospective birth cohort study Language: English Location: American	<u>Setting:</u> Healthy Start Study (n=1,063) conducted in the prenatal obstetrics clinics at University of Colorado Hospital in Aurora, Colorado. <u>Eligibility criteria:</u> Women (≥16 years) expecting a singleton birth, living in Colorado, and planning to deliver at University of Colorado Hospital. <u>Exclude criteria:</u> Women with serious chronic diseases (cancer, psychiatric diseases, steroiddependent asthma, pre-existent diabetes), as well as those who subsequently experienced a foetal death or delivered a severely premature infant (<32 week gestation) were excluded. <u>Sample size: n=804</u>	$\overline{\mathbf{x}} \pm \mathbf{SD}$ or \mathbf{n} (%) Age (year) 27.7 ± 6.1 Primiparous 287 (35.8) Pre-pregnancy BMI (kg/m²) 25.7 ± 6.3 Gestational length 39.4 ± 1.3 Fasting blood Yes	Enrolment time Gestational age at entry (≤24 gestation week) (All women were enrolled and delivered as of Nov 1, 2013) Length Follow up at least until birth Methods In-person research visits and hospital preconception visit Data collection Questionnaires, clinical diagnoses and medical records Loss to follow-up 0	median 27 week,	Birth weight was measured using a calibrated scale.	8
Hwang et al.2014	Study design: Prospective cohort study Language: English Location: Korea	Setting: The MOCEH study, a multicentre prospective hospital- and community-based cohort study in South Korea (n=1,751) Eligibility criteria: Pregnant women at mid-stage (15-28 gestation weeks) of a normal (not at risk) pregnancy who were willing to participate the MOCEH study. Exclude criteria: Twins (n=31), spontaneous abortion (n=23), intrauterine growth restriction (n=3), foetus congenital anomaly (n=12). Drop out (n=221), pregnancy complications (hypertension or/and diabetes, n=34). No information on dietary intake data	$\overline{\mathbf{x}} \pm \mathbf{SD}$ or \mathbf{n} (%) Age (year) 30.1 ± 3.6 Primiparous No statement Pre-pregnancy BMI (kg/m²) 21.3 ± 3.1 Gestational length 38.9 ± 1.4 Fasting blood No statement	Enrolment time Gestational age at entry (12-28 gestation week) (Aug 2006 to Dec 2010) Length Follow up until 5 years after delivery. Methods Clinical visits Data collection Questionnaires and medical records Loss to follow-up 221(17.94%)	Maternal serum <u>TG</u> was analysed twice at mid-pregnancy (12-28 gestational weeks) and at late pregnancy (29-42 gestational weeks) by means of an enzymatic method using an autonalyzer.		9

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		(n=135), total energy consumption <500 or >4000 kcal/day (n=5), No information on serum TG concentration at mid- or late pregnancy (n=276) $\underline{Sample\ size: n=1,011}$					
Kulkarni et al. 2013	Study design: Population-based birth cohort study Language: English Location: India	Setting: The Pune Maternal Nutrition Study (PMNS), a prospective birth cohort based on six rural villages in India. Eligibility criteria: Women with a singleton pregnancy of <21 weeks' gestation (n=797). Exclude criteria: Spontaneous abortions, fetal anomalies, multiple pregnancy, medical terminations late booking, Late abortions (n=12), late terminations (n=14), still birth (n=8), maternal death (n=1), congenital anomalies (n=9), baby not measured (n=51), mother diabetic (n=1), mother hypertensive (n=1), preterm (n=69) Sample size: n=631	$\overline{\mathbf{x}} \pm \mathbf{SD}$ or \mathbf{n} (%) Age (year) 21.4 ± 3.6 Primiparous 226 (35.8) Pre-pregnancy BMI (kg/m²) 18.0 ± 1.9 Gestational length 39.4 ± 1.7 Fasting blood Yes	Enrolment time Gestational age at entry (<21 gestation week) (June 1994 to April 1996) Length Follow up until birth. Methods No statement Data collection Questionnaires and clinical measurement Loss to follow-up 131 (16.44%)	Maternal fasting venous blood samples was collected at 18 and 28 weeks for total cholesterol HDL-C and triglycerides using standard enzymatic kits.	Measured by one of five trained fieldworkers within 72h of birth. <i>Birthweight:</i> measured by a Salter spring balance.	8
Vrijkotte et al.2012	Study design: Prospective cohort study Language: English Location: Netherlands	Setting: The Amsterdam Born Children and Their Development (ABCD) cohort study Eligibility criteria: Pregnant women visit to the obstetric care provider around the 12 th week of gestation agree to participant the ABCD biomarker study (n=4389) Exclude criteria: Women who had multiple gestation or who had no data on the gestational age at blood sampling, women with diabetes (preexistent as well as pregnancy induced), and	x ± SD or n (%) Age (year) 30.9 ± 4.9 Primiparous 2314 (57.7) Pre-pregnancy overweight or obese 830 (20.7) Gestational length No statement Fasting blood No.	Enrolment time Gestational age at entry (around 12 th gestation week) (Jan 2003 to Mar 2004) Length Follow up at least until birth. Methods Obstetric care provider visit and the Youth Health Care Registration and the Dutch Perinatal Registration (PRN).	during routine blood collection for laboratory TC and TG levels assessment during their first prenatal	Information on pregnancy outcomes was obtained from the Youth Health Care Registration and the Dutch Perinatal Registration (PRN). <u>SGA:</u> birth weight below the 10 th percentile for gestational age based on gender- and parity-specific standards from the PRN. <u>LGA:</u> birth weight above	8

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		those using lipid-altering medication (e.g. antiepileptic drugs, steroids, insulin, antidepressants, thyroid hormones, or sleep medication) were excluded. $\underline{Sample\ size\ :\ n=4,008}$		<u>Data collection</u> Questionnaires and Health care registration system. <u>Loss to follow-up</u> 381 (8.68%)		the 90 th percentile for gestational age based on the same gender0and parity-specific standards from the PRN.	
Retnakar en et al. 2012	Study design: Prospective cohort study Language: English Location: Canada	Setting: Ongoing prospective observational cohort study Eligibility criteria: White, Asian and South Asian pregnant women with term (37-41 weeks' gestation inclusive) singleton pregnancies were recruited at the second or early in the third trimester. Exclude criteria: Women with gestational diabetes. Sample size: n=472	$\overline{\mathbf{x}} \pm \mathbf{SD}$ or Median(IQR) Age (year) Lowest tertile birthweight: 33.6 \pm 4.0 Middle tertile birthweight: 34.5 \pm 4.3 Highest tertile birthweight: 33.6 \pm 4.0 Primiparous 251 (53.18) Pre-pregnancy BMI (kg/m²) Lowest: 22.6(20.7-25.4) Middle: 22.6(20.8-25.8) Highest: 23.6(22.3-27.4) Gestational length Lowest: 38.6 \pm 1.1 Middle: 39.2 \pm 1.0 Highest: 39.6 \pm 1.1 Fasting blood Yes.	Enrolment time Gestational age at entry (around 24 th -28 th gestation week) (No statement about recruitment time) Length Follow up until 3 months postpartum period Methods No statement Data collection No statement Loss to follow-up 0	Maternal fasting serum samples were obtained at the time of the oral glucose tolerance test (late second to early third trimester, median 30 week) for laboratory total cholesterol, HDL-c, LDL-c and triglycerides levels measurements.	Birthweight was measured at delivery. LGA: sex-specific birth weight for gestational age was above the 90 th percentile of Canadian foetal growth curves for the relevant ethnic group (white, Asian or South Asian) Macrosomia: birthweight over 4,000 g	7
Hou et al.2014	Study design: Prospective observational	Setting: Hospital-based study Eligibility criteria:	Median (25 th -75 th) <u>Age (year)</u> 26 (24-29)	Enrolment time Gestational age at entry (around 28 th – 37 th gestation	Maternal fasting venous blood was collected at the	LGA: birth weight were above the 90 th percentile for gestational age in	7

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
	study <u>Language:</u> English <u>Location:</u> China	Pregnant women with naturally conceive, singleton pregnancy during 28-37 week gestation were enrolled into this study <i>Exclude criteria:</i> Diabetes, abnormal glucose tolerance, chromosomal abnormality, inherited metabolic diseases thyroid disease, and risk for foetal chromosomal abnormality New-borns with preterm birth, inherited metabolic diseases, congenital abnormalities and congenital heart diseases. Sample size: n=2,790	Primiparous No statement Pre-pregnancy BMI (kg/m²) 19.93 (18.55- 21.63) Gestational length 39 (38-40) Fasting blood Yes.	week) (No statement about recruitment time) Length Follow up until delivery Methods Clinical visit Data collection Questionnaire, clinical measurement and diagnosis Loss to follow-up 0	enrolment time for laboratory TC, HDL-C,LDL-C and TG assay.	accordance with Neonatal Birth Weight for Gestational Age and Percentile in 15 cities in China.	
Kramer et al. 2014	Study design: Prospective cohort study Language: English Location: Canada	Setting: Ongoing prospective observational cohort study Eligibility criteria: Women with singleton delivery between April 2005 and January 2011, at term (≥37 weeks gestation, with infant birthweight >2500 g) Exclude criteria: No Sample size: n=340 (GDM, n=90; non-GDM, n=250)	x ± SD or n (%) Age (year) No statement Primiparous 340 (100) Pre-pregnancy BMI (kg/m²) No statement Gestational length No statement Fasting blood Yes.	Enrolment time Gestational age at entry (around 24 th -28 th gestation week) (Apr 2005 - Jan 2011) Length Follow up until 3-month postpartum period Methods Clinical investigation unit Data collection Questionnaire, clinical measurement Loss to follow-up 0	Maternal fasting serum samples were obtained at the time of the oral glucose tolerance test (late second to early third trimester, median 30 week) for laboratory total cholesterol, HDL-c and triglycerides levels measurements.	Infant weight gain at 3 months: the difference between weight at 3 months and birthweight. SD scores for weight gain at 3 months were determined for the study population, which was then stratified into two groups: infants weight rapid weight gain in the first 3 months (≥0.5 SD) and those without (<0.5 SD)	7
Harmon et al.2011	Study design: Prospective observational study Language: English Location:	Setting: Normal weight (BMI 20-25 kg/m²) and obese (BMI 30-38 kg/m²) women with NGT were enrolled at <15 weeks' gestation from the University of Colorado Hospital vicinity Eligibility criteria: Singleton pregnancies, being aged 18-35 years, being English speaking, and having a fasting blood glucose (FBG) <95 mg/dL.	31.2 ± 2.3 Obese: 26.5 ± 4.2 Parity Normal weight:	Enrolment time Gestational age at entry (<15 th gestation week) Length Follow up until birth. Methods No statement Data collection Questionnaire, clinical	Both early (14-16 weeks) and late (26-28 weeks) in gestation, all women had non-esterified free fatty acids (FFAs) measured. Triglycerides were	Birthweight.	6

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
	American	birthweight >2500 g) $\underline{Exclude\ criteria:}$ Having a history of diabetes, hypertension, triglycerides>300 mg/dL, chronic diseases; tobacco or alcohol use; or treatment with steroids/ β -blockers. Women with positive gestational diabetes diagnosis at baseline or 24-28 weeks' gestation were excluded. $\underline{Sample\ size: n=38}$	Pre-pregnancy BMI (kg/m^2) Normal weight: 22.4 ± 1.9 Obese: 33.1 ± 3.4 Gestational length Normal weight: 39.4 ± 0.3 Obese: 39.6 ± 0.3 Fasting blood No statement	measurement <u>Loss to follow-up</u> 4 (8.20%)	measured in early gestation only.		
Son et al.2010	Study design: Retrospective longitude observational study Language: English Location: Korea	Setting: No statement. Eligibility criteria: Pregnant women diagnosed with GDM by the OGTT with complete maternal overnight fasting blood samples within 2 weeks of GDM diagnosis. Exclude criteria: Women having hypertensive disorder (n=9), thyroid disorder (n=4), connective tissue disease (n=3). Patients who delivered before 35 weeks of gestation (n=14) and cases of foetal congenital malformation (n=10) or multifetal gestations (n=6) were also excluded. Sample size: n=104	38.3 ± 1.2	Enrolment time Gestational age at entry (24 th -30 th gestation week) Length Follow up until birth. Methods No statement Data collection clinical measurement Loss to follow-up 0	Maternal fasting serum TG, total cholesterol, low-density lipoprotein (LDL) and high-density lipoprotein (HDL) cholesterol concentrations at 24 th -32 th gestation week <i>Hypertriglyceridem ia</i> was defined as a TG level greater than the 75 th percentile value (<3.33 mmol/L)	Infants with birthweights above the 90th percentile were classified as LGA, based on gestational age and sex-adjusted birthweights from a Korean national database.	5
Ahmad. 2006	Study design: Controlled prospective study Language: English	Setting: Four antenatal clinics (ANC): Hospital Universiti Sains Malaysia, Kota Bharu Health Cinic, Kubang Kerian Health Clinic and Kedai Lalat Health Clinic. Eligibility criteria: Pregnant women attending the antenatal clinics at gestation between 24 to 32 weeks	$\overline{\mathbf{x}} \pm \mathbf{SD}$ $\underline{Age (year)}$ 30.87 ± 6.70 $\underline{Gravidity}$ 3.76 ± 2.69 $\underline{BMI (kg/m^2)}$ 23.36 ± 4.04 $\underline{Gestational \ length}$	Enrolment time Gestational age at entry (24 th -32 th gestation week) Length Follow up until delivery. Methods Antenatal clinics visit and appointment	Maternal fasting lipid profile was taken at between 24 to 32 weeks gestation for laboratory analyses. (total cholesterol and	At delivery, weight of the newborn were noted. LGA: Neonatal birth weight above the 90 th percentile of gender specific birth weight curve of Malaysia.	7

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
	Location: Malaysia	gestation. Exclude criteria: Diabetic (diagnosed diabetic prior to conception and gestational diabetes requiring insulin); Hypertension or preeclampsia (hypertensive disorder), lupus and antiphospholipid syndrome, fetal anomaly diagnosed through ultrasound during booking or noted abnormal at birth; multiple gestation; pre-term delivery. Sample size: n=246	39.00 ± 1.29 <u>Fasting blood</u> Yes.	Data collection clinical records Loss to follow-up 50 (13.9%)	triglycerides)		
Di et al.2005	Study design: prospective observational study Language: English Location: Italy	Setting: The diabetes Section of the Department of Endocrinology and Metabolism of the University of Pisa, Italy. Eligibility criteria: Pregnant Caucasian women with positive diabetic screening performed at 24 to 30th week of gestation, Exclude criteria: Women with hypertensive disorders, thyroid disorder, lupus and antiphospholipid syndrome. Sample size: n=180 (NGT=121) The main analysis of our interest is conducted on NGT women who delivered at term. (n=83)	$\overline{\mathbf{x} \pm \mathbf{SD}}$ or \mathbf{n} (%) $\frac{Age (year)}{33 \pm 4}$ $\frac{Primiparous}{106 (59)}$ $\frac{Pre-pregnancy}{BMI (kg/m^2)}$ 23.6 ± 4 $\frac{Gestational \ length}{39.3(39-40)}$ $\frac{Fasting \ blood}{Yes}$	Enrolment time Gestational age at entry (24 th -28 th gestation week) Length Follow up until delivery. Methods Antenatal clinics visit and appointment Data collection clinical records Loss to follow-up 0	Maternal overnight fasting lipid level (Total cholesterol, LDL-C, HDL-C, Triglycerides) at between 24 th and 28 th week of gestation.	Birthweight. Macrosomia: neonatal body weight over 4kg or as a neonatal weight greater than 90 th percentile for gestational age (LGA), according to the reference table.	5
Schaefer -Graf et al.2008	Study design: Secondary analysis of RCT study Language: English Location:	Setting: Two hospital based diabetic prenatal care clinics. Original study (n=199): Women diagnosed as GDM based on a 75-g OGTT in capillary blood. (capillary fasting glucose <120 mg/dl, postprandial glucose <200 mg/dl). This analysis (n=150):	31.2 ± 4.9 \underline{Parity} 2.05 ± 1.2 $\underline{Pre-pregnancy}$	Enrolment time Gestational age at entry $(28.3 \pm 2.4 \text{ weeks});$ $(Jan 2000 - Jan 2003)$ Length Follow up until day 2 after delivery Methods Clinical visits (28, 32, 36,	FFAs, cholesterol and triglycerides were measured every clinical visit (28, 32, 36 and close to delivery) using commercial		5

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
	German	Accepted insulin therapy; availability of complete maternal blood and cord blood samples.		39 weeks, labour and day 2 postpartum) <u>Data collection</u> No statement <u>Loss to follow-up</u> 49/199 (24.6 %)		classified as SGA, and those with birth weight > 90 th percentile as LGA based on gestational age and sexadjusted birth weight percentiles derived from a German national database. Cord blood samples ware taken immediately following delivery and serum was stored at -80°C for TGs, free fatty acids(FFAs) and cholesterol measurements.	
Swierze wska et al. 2015	Study design: Prospective observational study Language: English Location: Poland	Setting: No statement Eligibility criteria: 136 Caucasian women were included into this study: 106 diagnosed with GDM and 31 pregnant women with normal glucose tolerance. Exclude criteria: No statement Sample size:136 GDM group: 106 NGT group: 31	x ± SD or n (%) Age (year) GDM: 30.2±0.36 NGT: 28.87±0.6 Primiparous No statement. Pre-pregnancy weight (kg) GDM:25.29±0.4 NGT: 23.05±0.52 Gestational length (days) No statement Fasting blood No statement.	Enrolment time Gestational age at entry (No statement); (2012 - 2013) Length Follow up until birth. Methods No statement Data collection Survey, interview Loss to follow-up 0	blood samples were collected twice (27-32 wks	Macrosomia was diagnosed in newborn with the firth weight of ≥4000 g, and LGA if the birth weight exceeded the 90 th percentile.	5
Sommer et al.2015	Study design: Population- based, multi- ethnic,	<u>Setting:</u> The STORK Groruddalen study (n=823), a population-based cohort study of healthy pregnant women attending Child Health	29.3 ± 4.8	Enrolment time Gestational age at entry (<20 gestation week) In practice, the STORK	total-, HDL- and LDL-cholesterol	Birth weight was measured with calibrated electronic scales immediately after birth.	9

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
	prospective cohort Language: English Location: Norway		319 (45.6) <u>Pre-pregnancy</u> <u>BMI (kg/m²)</u> 24.6 ± 4.8 <u>Gestational length</u> (<u>days)</u> 281 ± 9 <u>Fasting blood</u> Yes.	study also includes 77 (9.4 %) and 11 (1.3%) women entry into this study at 20-24 gestation week and later than gestational week 24, respectively. (May 2008 to May 2010) Length Follow up at least until 3 days after birth. Methods Clinic visits Data collection Questionnaires, clinical measurement and laboratory diagnosis. Loss to follow-up 37(5.29%)	from venous blood with a colorimetric method at the	skinfolds were measured to the nearest 0.2mm with a skinfold calliper	
Slagjana et al.2014	Study design: Population- based, multi- ethnic, prospective cohort Language: English Location: Norway	Setting: The Outpatient Department of the University Endocrinology, Diabetes and Metabolic Disorders Clinic Eligibility criteria: GDM women with singleton pregnancy, and the neonates were delivered at the University Gynaecology and Obstetrics Clinic. Exclude criteria: None Sample size: n=200	x ± SD or n (%) Age (year) LGA: 31.4±5.6 AGA: 31.1±5.6 SGA: 32.9±5.1 Primiparous No statement Pre-pregnancy BMI (kg/m²) LGA: 28.4±6.1 AGA: 26.5±4.9 SGA: 25.0±4.6	Enrolment time No statement on recruitment date and entry gestational age. Length Follow up until birth. Methods Clinic visits Data collection Clinical measurement and laboratory diagnosis. Loss to follow-up	Maternal overnight fasting blood samples were collected at the second half of pregnancy(LGA; 28.6±7.7; AGA: 28.0±7.1; SGA: 23.8±7.6) for Total cholesterol, HDL-C,LDL-C and triglycerides	<u>LGA:</u> birth weight above the 90 th percentile. <u>SGA:</u> birth weight below the 10 th percentile for gestational age. <u>AGA:</u> birthweight between LGA and SGA.	5

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
			Gestational length (weeks) LGA: 39.3±1.5 AGA: 38.2±1.9 SGA: 36.4±3.7 Fasting blood Yes.	0	laboratory assessment.		
	Study design: Prospective cohort Language: English Location: Iran	Setting: Shariati Hospital affiliated to Tehran University of Medical Sciences, Tehran, Iran Eligibility criteria: Pregnant women were diagnosed with GDM. Exclude criteria: Women with a history of systemic underlying diseases (cardiovascular, renal, thyroid, liver, autoimmune and connective tissue disorder), substance abuser, overt diabetes mellitus (except previous history of GDM), multifetal gestations and major fetal malformation. Sample size: n=112	x ± SD or n (%) Age (year) 27.23±4.19 Parity 2.74 (66.1) Pre-pregnancy weight (kg²) 67.40±10.00 Gestational length (days) No statement Fasting blood Yes.	Enrolment time Gestational age at entry (27.02 ± 0.68 weeks); (Mar 2011 - May 2012) Length Follow up until birth. Methods Clinic visits Data collection A combination of interviews and questionnaires in timing of glycemic screening (24-28 weeks) Loss to follow-up 20 (15.15%)		SGA: birthweight <10 th percentile. LGA: birthweight >90 th percentile. Macrosomia: >4000 g	7
	Study design: Prospective cohort Language: English Location: Ireland	Setting: The Perinatal day centre of University Maternity practice. Eligibility criteria: White European women with an ongoing singleton pregnancy were enrolled when they were referred to the Perinatal day centre for OGTT screening test. Exclude criteria: Women who were unable to give informed consent or who were less than 18 years of age were excluded.	x ± SD or n (%) Age (year) 32±5 Primigravidas 67(35.4) Pre-pregnancy BMI (kg/m²) No statement Gestational length (days) 277±14 Fasting blood	Enrolment time Gestational age at entry (when women attend OGTT screening test); (Mar 2011) Length Follow up until birth. Methods Clinic visits Data collection Clinical measurements, diagnosis, hospital's	Maternal fasting venous blood sample was obtained to measure the TC, HDL-C, LDL-C and TG when women attend OGTT screening test.	After delivery, birthweight was obtained from the Hospital's computerized database.	5

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		Sample size: n=189	Yes.	computerized database. <u>Loss to follow-up</u> 0			
Zhou et al.2012	Study design: Prospective cohort Language: English Location: China	Setting: Routine obstetric care in the Nanjing drum tower hospital Eligibility criteria: Nulliparous pregnant women < 20 weeks gestation visited the antenatal department and had booked to deliver their infants at Nanjing Drum Tower Hospital. Exclude criteria: Women with family history of dyslipidemia, chronic diseases that may affect the lipid profile such as hypertension, diabetes mellitus and systemic lupus erythematosus, or used a medication that affected the lipid profile. Sample size: n=1,000	x ± SD or n (%) Age (year) 28.6±3.4 BMI (kg/m²) 22.54±2.86 Gestational length (weeks) 39.3±1.2 Fasting blood Yes.	Enrolment time Gestational age at entry (20 gestation week); (Jun 2009 to Jan 2010) Length Follow up until birth. Methods Clinic visits Data collection Clinical measurement and laboratory diagnosis Loss to follow-up 15 (1.5%)	Maternal overnight fasting blood at 20 weeks gestation were measured for serum TG, TC, LDL-c and HDL-c. Hypo-HDL-cholesterolemia was defined as fasting serum HDL-C levels below the optimal cut-off value.	Infants with birthweight <10 th percentile were classified as SGA based on gestational age and sex adjusted birth weight percentiles, and those with birth weight above 4,000 g were classified as macrosomia.	5
Vrijkotte 2011	Study design: Prospective community- based cohort study Language: English Location: Netherlands	Setting: Amsterdam Born Children and their Development (ABCD) study Eligibility criteria: All pregnant women living in Amsterdam were invited to enrol in the ABCD study at their first prenatal visit to an obstetric care provider at about the 12 th week of gestation. Exclude criteria: Women who gave birth to twins, delivered preterm (<37 wks), with known diabetes (pre-existent as well as pregnancy related), or whose infants had congenital abnormalities were excluded. Women who used lipid-altering medication, such as antiepileptic drugs, steroids, insulin, antidepressants, thyroid hormones, or sleep	$\overline{\mathbf{x} \pm \mathbf{SD}}$ or n (%) Age (year) 31.0±4.8 Pre-pregnancy BMI (kg/m²) <18.5: 115(4.6%) 18.5-24.9: 1869(74.7%) 25.0-29.9: 388(15.5%) ≥30: 130(5.2%) Primigravidas 1412(56.4) Gestational length (weeks) 37-40 wks: 1779(71.6%)	Enrolment time Gestational age at entry (around 12 gestation week); (Jan 2003 to Mar 2004) Length Follow up until 12 months after birth. Methods Clinic visits Data collection Clinical measurement and laboratory diagnosis Loss to follow-up 0	Maternal non- fasting serum samples were taken during routine blood collection for screening purposes after the first prenatal check-up for lipid laboratory measurements (TG and TC).	Birthweight for gestational age SDS was determined based on sex-and partiy-specific standards from the Dutch Perinatal Registry. In the first year, weight and length were measured on average 8 times. Weight, length and BMI were expressed as SDS by using internal sexspecific reference curve from the ABCD study. To further explore postnatal growth, the	7

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		medication also were excluded. Sample size: $n=2,502$	4143 wks: 707(28.4%) <i>Fasting blood</i> No.			amount of accelerated growth was defined as an increase >0.67 SDS between 2 time points (between 1 and 6 months of age)	
Vinod et al. 2011	Study design: Ongoing prospective cohort study Language: English Location: American	Setting: University of Michigan Health System Eligibility criteria: Eligible participants were 18-45 years of age, between 6 and 10 weeks gestation with a singleton pregnancy, and intended to deliver at the study hospital. Exclude criteria: Participants who did not complete the study and delivered a live infant. 1% of women were excluded from any analysis because of missing data. Sample size: n=143	$\overline{\mathbf{x}}$ ± SD or n (%) Age (year) ≤30: 79(55.2) >30: 64(44.8) <u>Pre-pregnancy</u> <u>BMI (kg/m²)</u> Normal weight: 72 (50.4) Overweight/Obese: 71 (49.6) <u>Primigravidas</u> 54 (37.8) <u>Gestational length</u> (days) 274.0 ± 13.2 <u>Fasting blood</u> No.	Enrolment time Gestational age at entry (6-10 gestation week); (No statement on entry date) Length Follow up until birth. Methods Clinic visits Data collection Interview, Questionnaire, Medical records, Clinical measurement and laboratory diagnosis Loss to follow-up (1%)	time points during pregnancy: 6-10, 10-14, 16-20, 22- 26 and 32-36 weeks gestation for laboratory lipid measurements (TC,	Infant birthweight was collected at delivery. The residual values from each fit were used to represent the gestational age-adjusted birthweight (aBW).	6
	Study design: prospective observational study Language: English Location: Poland	Setting: Department of Obstetrics and Women Diseases for a tertiary-level, specialistic antenatal care. Eligibility criteria: GDM diagnosed following WHO criteria, singleton pregnancy, live birth and no fetal malformation suspected during gestation or detected postpartum. Exclude criteria: None. Sample size: n=357	Median (min-max) Age (year) 29 (17-48) Pre-pregnancy BMI (kg/m²) 24.2 (16.7-46.1) Primigravidas No statement Gestational length (weeks) 38 (32-42) Fasting blood	Enrolment time Gestational age at entry (GDM diagnosis week); (1993 to 2005) Length Follow up until birth. Methods Clinic visits Data collection Clinical measurement and laboratory diagnosis Loss to follow-up 0	Maternal overnight fasting blood sample were taken for laboratory lipid assessment (TC, HDL and triglycerides) at their first booking weeks (GDM diagnosis week)	Birth weight and the proportion of LGA (defined as a birth weight >90 th percentile for local population after adjusting for gestational age and sex) was studied at the end-point.	5

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
Clausen et al.2005	Study design: Prospective cohort study Language: English	Setting: Aker Hospital in the Oslo city area Eligibility criteria: All pregnant women living in Oslo area were offered an ultrasound investigation at 17-19 weeks of gestation	Yes. $\overline{\mathbf{x} \pm \mathbf{SD}}$ or \mathbf{n} (%) \underline{Age} (year) 29.9±4.4 \underline{The} 1 st trimester \underline{BMI} (kg/m²) 23.0±3.7	Enrolment time Gestational age at entry (17-19 th gestation week); (1995-1996) Length Follow up until birth.	Maternal fasting blood samples were drawn at 17- 19 th gestation weeks for laboratory lipid	Macrosomia: birth weight above 4,500 g or a z-score above the 95 percentiles.	7
	<u>Location:</u> Norway	Exclude criteria: Pre-gestational diabetes, multiple pregnancies, preterm births, missing medical records, no information on birth weight, lost for follow-up Sample size: n=2,050	Primigravidas 1030(50.3) Gestational length (weeks) 39.7±1.3 Fasting blood Yes	Methods Clinic visits Data collection Clinical measurement and laboratory diagnosis Loss to follow-up 244(10.6%)	measurements (TGs, TC, HDL-C, non-HDL- cholesterol).		
Mathews et al.2003	Study design: Prospective cohort study Language: English Location: United Kingdom	Setting: The geographic catchment area of St Mary's Hospital, Portsmouth, United Kingdom Eligibility criteria: White nulliparous women attending their first hospital antenatal clinic were stratified by self-reported smoking status. Simple random selection was carried out within each stratum. Exclude criteria: Preterm birth, insufficient blood for assays and still birth Sample size: Subjects for birth weight and early pregnancy nutrition analyses: n=798	$\overline{x} \pm SD$ or n (%) Age (year) 25.4±4.9 Pre-pregnancy BMI (kg/m²) 23.1±3.9 Gestational length (days) Boys: 280.3±9.9 Girls: 281.3±9.5 Fasting blood NS	Enrolment time Gestational age at entry (14-17 th gestation week, range: 9-20 wk); (May 1994 – Feb 1996) Length Follow up until birth Methods Clinical visits Data collection Questionnaire, Clinic measurement and laboratory diagonosis Loss to follow-up 0	Maternal blood samples were obtained from subjects at two time points (early pregnancy: at around 16 gestation week, later pregnancy: at around 28 gestation week) for total cholesterol laboratory analyses	Infants were weighed at delivery to the nearest 5 g on digital scales.	8
Olmos et al.2014	Study design: Prospective observational study Language: English	Setting: Obstetricians Eligibility criteria: Women aged 18-42 years with singleton pregnancy, under the care of an Obstetrician of the University Health Care Network, having GDM confirmed recently (<14 days)	32.7±5.3	Enrolment time Gestational age at entry (after GDM diagnosis week); (Jan 2009 – Jun 2013) Length Follow up until birth	Maternal fasting lipid (triglycerides, total cholesterol, HDL-C) level were measured in the 2 nd and 3 rd trimesters. All lipid	Birth weight z-scores. Macrosomia: a birth weight above 90 th percentile, was used, applying to that effect the tables of the Chilean	6

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
	Location: Chile	by an oral glucose tolerance test (OGTT) test. Exclude criteria: Women unable to give informed consent or who were less than 18 years of age were excluded. Sample size: n=279 Normal weight group: n=128 Overweight group: n=105 Obese group: n=46	32.3±4.7 <u>Primiparous</u> No statement <u>Pre-pregnancy</u> <u>BMI (kg/m²)</u> Normal weight: 22.3±1.5 Overweight: 26.1±3.1 Obese: 33.1±2.7 <u>Gestational length</u> (weeks) Normal weight: 38.0±1.3 Overweight: 37.7±1.7 Obese: 37.6±1.7 <u>Fasting blood</u> Yes.	Methods Clinic visits Data collection Clinical measurements and diagnosis, and laboratory diagnosis Loss to follow-up 0	parameters were calculated as z-scores based on Alvarez paper.	Ministry of Health, in use since 2004.	
Emet et al.2013	Study design: Prospective observational study Language: English Location: Turkey	Setting: Antenatal care, Eligibility criteria: 1,000 pregnant patients between 17 and 48 years of age were included in this prospective longitudinal and uni-centre study. Exclude criteria: Patients with type I-II diabetes mellitus and hypothyroidism, multiple gestations, dyslipoproteinemia were excluded from the study. Also, patients on special diets because of underlying diseases or personal preferences such as gluten or casein-free diets, vegetarian diet, liver or renal failure diet, etc., or patients using medications that effect lipid metabolism were excluded as well. Patients whose pregnancies were	x ± SD or n (%) Age (year) 28.5±5.5 Parity 0.94±0.98 Pre-pregnancy BMI (kg/m²) No statement Gestational length (weeks) 38.9±1.8 Fasting blood Yes	Enrolment time Gestational age at entry (<14 gestation week); (Jan 2010 – Dec 2011) Length Follow up until birth Methods Clinic visits Data collection Questionnaire, interview, clinical and laboratory diagnosis Loss to follow-up 76(8.68%)	Maternal lipid profile (TG, TC, HDL, LDL) were tested at the first antenatal visit (<14 weeks) and the last trimester (>28 weeks)	Birthweight was recorded. Third month infant weight was also surveyed.	5

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		terminated before 24 gestational week, patients who dropped out of routine antenatal and patients who gave birth outside the hospital were also not included in this analysis Sample size: n=801					
Liu et al. 2016	Study design: Retrospective cohort study Language: English Location: China	Setting: The first affiliated hospital of Sun Yat-sen University Eligibility criteria: Singleton pregnant women who underwent a FPG test at the first prenatal care, and delivered in our centre were recruited for the present study. Exclude criteria: Pregnant women with overt DM before, pregnancy or treated with insulin during gestation were excluded in the present study Sample size: n=1,546	Age (year) GDM: 31.85±4.24 NGT: 29.42±3.82 Primiparous GDM: 234 (84.7) NGT: 969 (76.2) Pre-pregnancy BMI (kg/m²) GDM: 21.20±3.00 NGT: 20.47±2.60 Gestational length (days) GDM: 271.33±11.70 NGT: 273.94±11.91 Fasting blood YES.	Enrolment time Gestational age at entry (10 th -24 th gestation week); (Jan - Dec 2013) Length Follow up until birth Methods Clinic visit Data collection Questionnaire, clinical measurements and diagnosis, laboratory diagnosis. Loss to follow-up 0	Maternal fasting venous plasma were obtained at the first prenatal visit (24-28 gestational weeks) for the examination of lipid profiles (triglyceride, cholesterol, LDL, HDL)	Neonatal birth weight was measured with a calibrated electronic scale.	7
Brunner et al. 2013	Study design: Secondary analyses of RCT study Language: English Location: German	Setting: The Impact of Nutritional Fatty Acid on Infant Adipose Development (INFAT) study, an open-label randomized controlled trial Eligibility criteria: Healthy pregnant women with singleton pregnancies and a pre-pregnancy BMI between 18 and 30 kg/m² were enrolled and randomly assigned to either an intervention (n=104) or a control group (n=104) from the	x ± SD or n (%) <u>Age (year)</u> 31.8±4.7 <u>Primiparous</u> 122(58.5) <u>Pre-pregnancy</u> <u>BMI (kg/m²)</u> 22.3±3.0 <u>Gestational length</u> (weeks) 39.6±1.5	Enrolment time Gestational age at entry (before 15 th gestation week); (No statement on recruitment date) Length Follow up until 2 years old. Methods Clinic visits Data collection Clinic measurement,		The infants were examined at birth (for skinfolds: 3-5 days post-partum), at 6 weeks, 4months, 1 and 2 years post-partum. Birthweight was retrieved from the medical record. Anthropometric measurements of the	7

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		15 th week of gestation until 4 months post- partum. <i>Exclude criteria:</i> None. <i>Sample size : n</i> =208	Fasting blood YES	medical records, clinic diagnosis, laboratory analyses <u>Loss to follow-up</u> 0		infants were taken by trained investigators according to standardized procedures. Skinfolds were measured in triplicate with a Holtain calliper at the left body axis at four sites (triceps, biceps, subscapular and suprailiac).	
Knopp et al.1992	Study design: Prospective observational study Language: English Location: American	Obstetrical practices at the two main Group		Enrolment time Gestational age at entry (24 th – 32 nd gestation week); (Jan 1985 – May 1986) Length Follow up until birth Methods Clinic visit Data collection Medical records, laboratory measurement. Loss to follow-up 0	Maternal overnight fasting blood samples collected at between 24 th and 32 nd gestation was measured by laboratory for plasma triglycerides.	Birthweight was adjusted for differences in gestational age by dividing the observed birth weight by the 50 th percentile birth weight for that gestational age, giving a birth-weight ratio.	6

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		Positive screenees(PS+): n=264 GDM: n=96					
Knopp et al.1985	Study design: Prospective observational study Language: English Location: American	Setting: Group Health Cooperative of Puget Sound, a prepaid health program. Eligibility criteria: Subjects were identified at 26-28 wk gestation by a prospective random sampling scheme, were invited to participate, and, after consent was given, had anthropomorphic measurements and blood sampled at home at 36 wk gestation by a visiting research nurse. Exclude criteria: Women were excluded if they aborted or delivered before 36 wk or had fasted <12 h. women who were not Caucasian, were under 18 yr of age, or had a twin pregnancy were also excluded. Sample size: n=283	x ± SD or n (%) Age (year) 28.0±3.8 Primiparous 102 (36) Pre-pregnancy BMI (kg/m²) No statement Gestational length (days) 283.4±18.6 Fasting blood Yes	Enrolment time Gestational age at entry (26-28 gestation week); (No statement on recruitment date) Length Follow up until birth. Methods Clinic visit, home visit Data collection Interview, hospital records, clinical and laboratory measurements. Loss to follow-up 10 (3.5%)	Maternal fasting blood sampled at home at 36 wk gestation by a visiting research nurse for laboratory lipid measurements (HDL-C, VLDL-C, LDL-C and FFA)	Birth weight data were extracted from hospital records. Birth weight was adjusted for gestational age and expressed as the birth weight ratio as determined from the expected date of confinement by dividing the observed birth weight by the median expected for gestational age using the University of Oregon (sea level) tables.	7
	Study design: Prospective observational study Language: English Location: German	Setting: Vivantes Medical Center Department of Obstetrics in Berlin Eligibility criteria: 1)documented normal 75-g oral glucose tolerance test according to Carpenter and Coustan criteria (5.0/10.0/8.6 mmol/L) with three glucose values in capillary blood using the hexokinase method; 2) accurate gestational age, confirmed by an ultrasound examination before 20 weeks of gestation; 3) singleton pregnancy; 4) absence of identified fetal anomalies; 5) delivery after 34 weeks; 6) signed informed consent Exclude criteria: No statement	x ± SD or n (%) Age (year) 30.0±0.4 Parity 2.07±0.09 Pre-pregnancy BMI (kg/m²) 25.7±0.4 Gestational length 38.8±0.1 Fasting blood Yes	Enrolment time Gestational age at entry (No statement on recruitment gestation week); (Aug 2007 – Aug 2008) Length Follow-up until 48h after birth Methods Hospital stay Data collection Laboratory diagnosis. No statement around how did they get maternal baseline information. Loss to follow-up	Maternal overnight fast blood samples were taken from a radial vein either on the morning of admission for surgery in cases of primary Caesarean section or at the last visit o the obstetrical clinic, no longer than 1 week before delivery. Serum triacylglycerols, free fatty acids and	obtained shortly after delivery and neonatal skinfold thickness at the flank was measured within 48 h to calculate fat mass. LGA: birthweight <10 th percentile. SGA: birthweight >90 th percentile. Cord blood samples from one of the umbilical arteries were taken immediately after	5

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		Sample size: n=190		0	cholesterol were measured in laboratory.	Serum glucose, insulin, triacyglycerols, free fatty acids and cholesterol were measured in cord blood.	
Nolan et al.1995	Study design: Prospective observational study Language: English Location: Australia	Setting: Obstetric clinic at the Mercy Hospital for Women Eligibility criteria: Women with singleton pregnancies had routine 3 rd -trimester oral glucose tolerance tests performed and have been included for analyses in this study. Exclude criteria: No statement Sample size: n=388	x ± SD or n (%) Age (year) 28.4±5.3 Primiparous No statement BMI at wk 20 (kg/m²) 24.7±4.2 Gestational length No statement Fasting blood No.	Enrolment time Gestational age at entry (≤20th gestation week); (1991) Length Follow up until birth Methods clinic visits Data collection Clinic records, clinic visits, laboratory measurements Loss to follow-up 0	During the morning of the first clinic visit (average sampling time: 12.2±6.2 weeks), all women had non-fasting serum TG and cholesterol measured within their routine antenatal screening blood analyses. TG and cholesterol were assayed by enzymatic colorimetric methods.	(BWR) for all infants was calculated by dividing the observed birth weight by the 50 th percentile birth weight	6
Friis et al.2012	Study design: Prospective observational study Language: English Location: Norway	Setting: A subcohort of the STORK study, Eligibility criteria: women of Scandinavian heritage (n= 1031) who registered for obstetric care at Oslo University Hospital - Rikshospitalet Exclude criteria: Multiple pregnancies, known pregestational diabetes, and severe chronic diseases (lung, cardiac, gastrointestinal or renal). Sample size: n=207	x ± SD or n (%) Age (year) 31±3.5 Primiparous 91(44) Pre-pregnancy height(cm)/weight (kg²) 168/66 Gestational length 40.1±1.4 Fasting blood Yes	Enrolment time Gestational age at entry (14 th -16 th gestation week); (2001-2008) Length Follow up until 4 days postpartum Methods Clinic visits Data collection Interview, clinic measurements, hospital records Loss to follow-up	Maternal fasting blood samples were collected at 30-32 th gestation weeks for total cholesterol, HDL, triglycerides, free fatty acids laboratory measurements.	Birthweight	6

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
				0			
Lei et al.2016	Study design: Prospective cohort study Language: English Location: China	Setting: The Department of Obstetrics of Guangdong Women and Children Hospital, Guangzhou, Guangdong Province Eligibility criteria: Pregnant women were recruited before 20 gestation wks Exclude criteria: Multiple pregnancy, conception by means of gonadotropin ovulation induction or in vitro fertilization, ischemic heart disease, stroke, peripheral vascular disease, dyslipidaemia, diagnosis of diabetes or/and hypertension before the current to participate in the study. Sample size: n=5,535	x ± SD or n (%) Age (year) 29.07±5.04 Primiparous 3152 (56.95) Pre-pregnancy BMI (kg/m²) 20.87±2.81 Gestational length 38.20±2.81 Fasting blood Yes.	Enrolment time Gestational age at entry (<20 th gestation week); (Jan 2012 – Dec 2014) Length Follow up until birth Methods Clinic visits Data collection Laboratory assessment, medical surveillance. Loss to follow-up 485 (8.06%)	Maternal fasting venous blood samples were drawn before 20 weeks to assess metabolic profile (TG and HLD-C). High level of TG was defined as ≥3.49 mmol/L (≥75 th percentile). Low level of HDL-C was defined as <1.3 mmol/L (<25 th percentile)	A newborn was considered SGA or LGA if birth weight as smaller or greater than the estimated 10 th /90 th percentile for the baby's gender and gestational age according to the Chinese data published before.	6
Kitajima et al.2001	Study design: Prospective observational study Language: English Location: Japan	Nagasaki University Hospital Eligibility criteria: Japanese pregnant women who had positive diabetic screen test results (at least 135mg/dl of plasma glucose level at 1 hour after 50-g oral glucose challenge) and a normal 75-g oral GTT. Exclude criteria: Women with pregestational or gestational diabetes mellitus were excluded. We also excluded women with hypertensive disorder, thyroid disorder, lupus, and antiphospholipid syndrome. Subjects who delivered before 37 weeks' gestation and cases of foetal congenital malformation or multifetal gestation were also excluded. Sample size: n=146	x ± SD or n (%) Age (year) 32±4 Primiparous 65(44%) Pre-pregnancy BMI (kg/m²) 21.2±2.7 Gestational length 39.0±1.2 Fasting blood Yes.	Enrolment time Gestational age at entry (24-32 gestation week); (Nov 1992 and Oct 1999) Length Follow up until delivery. Methods Clinic visits Data collection Self-report, clinic measurements and diagnosis Loss to follow-up 0	Maternal fasting blood samples were drawn to measure serum triglyceride, free fatty acids and total cholesterol levels at 24-32 gestation week through laboratory measurements. Maternal hyperlipidaemia was defined as a value higher than the 75th percentile value of each lipid concentration.	Neonatal birth weight above the 90th percentile of the gender specific Japanese birth weight curve was defined as <i>LGA</i> .	6
M	Study design:	Setting:	$\bar{x} \pm SD$	Enrolment time	Maternal blood	Macrosomia was defined	5

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
bi et al. 2014	Cohort study <u>Language:</u> English <u>Location:</u> Iran	The prenatal clinic of the Shahid Akbar Abadi Hospital <i>Eligibility criteria:</i> All women were generally healthy pregnant women carrying a single foetus, between 25 weeks and 32 weeks of their gestational age, BMI between 17.5 kg/m2 and 29 kg/m,2 without a history of diabetes prior to or during previous pregnancies and with a negative result from the diabetes screening test in the current pregnancy, hypertensive disease and preeclampsia, thyroid diseases, lupus, antiphospholipid antibody syndrome, and other collagen vascular diseases. <i>Exclude criteria:</i> Exclusion criteria were preterm labour prior to 37 weeks of gestational age and any abnormality or disorder in the foetus or neonate. <i>Sample size: n=154</i>	Age (year) 26.6±5.17 Parity 1.7±0.79 Pre-pregnancy BMI (kg/m²) 22.6±2.3 Gestational length No statement Fasting blood Yes.	Gestational age at entry (25-32 th gestation week); (2010-2011) Length Follow up until birth Methods Clinic visits Data collection Clinic measurement and diagnosis. Laboratory measurements. Loss to follow-up 16 (8%)	sample for checking fasting triglyceride (TG), total cholesterol, low-density lipoprotein (LDL), and high-density lipoprotein (HDL) after 10-12 hours of fasting at 25-32 th gestation week. (gestational age at the time of blood sampling: 30±2.1)	as neonate birth weight higher than 4000 g. LGA was defined as neonate's birth weight higher than 3412 g for infants at 38 weeks of gestational age, 3622 g for infants at 39 weeks of gestational age, 3798 g for infants at 40 weeks of gestational age, and 3930 g for infants at 41 weeks of gestational age. This definition was according to the neonates' weight higher than 75% of their predicted value according to their gestational age.	
	Study design: Secondary analyses of RCT study Language: English Location: Ireland	Setting: Randomised cOntrol trial of Low glycaemic index diet vs no dietary intervention in pregnancy to prevent recurrence of a large baby (ROLO) study, which was carried out in The National Maternity Hospital, Dublin, Ireland. Original study: Eight hundred secundigravida women who did not have gestational diabetes but had previously given birth to a macrosomic baby (birth weight equal to or above 4.0 kg), and were therefore at increased risk of delivering another macrosomic infant, were randomised to receive low glycaemic index (GI) dietary advice or usual antenatal care,	x ± SD or n (%) Age (year) 33.10±3.90 BMI at 14 weeks' gestation(kg/m²) 26.40±4.60 Gestational length (days) 282.80±7.50 Fasting blood Yes	Enrolment time Gestational age at entry (<14 th gestation week); (No statement on recruitment time) Length Follow up until 2 years old. Methods Clinic visits and follow-up appointments Data collection Clinic measurements, laboratory measurements. Loss to follow-up 0	Maternal fasting blood samples were taken in early pregnancy (approximately 14 th gestation weeks) and late pregnancy (28 th gestation weeks) for serum total cholesterol, HDL-C and triglyceride laboratory measurements. LDL-C concentration was	Infants were measured at birth, 6 months and 2 years of age for weight and recumbent length along with abdominal circumference and bicep, tricep, subscapular and thigh skinfold thicknesses.	7

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		which did not include dietary advice. Eligibility criteria: No statement. Exclude criteria: No statement. Sample size: n=331			estimated using the Friedewald equation.		
Jin et al.2016	Study design: Cohort study Language: English Location: China	Setting: Women's Hospital, Zhejiang University School of Medicine Eligibility criteria: 1) pregnant at 28–37 gestational weeks; 2) had integrated medical records and clear gestational age; 3) singleton pregnancy; and 4) naturally conceived. Inclusion criteria for newborns were singleton and 5-min-postpartum Apgar scores ≥ 7. Exclude criteria: 1) multiple pregnancy; 2) had diabetes mellitus, chromosomal abnormalities, inherited metabolic diseases or thyroid diseases before pregnancy; 3) experienced serious infection during early pregnancy; and 4) conceived with assisted reproductive techniques. Exclusion criteria for newborns were chromosomal abnormalities, inherited metabolic diseases and congenital abnormalities. Sample size: n=934	x ± SD or n (%) Age (year) 29.21±3.76 Primiparous 778(83.3%) Pre-pregnancy BMI (kg/m²) 20.66±2.70 Gestational length 38.84±1.22 Fasting blood Yes.	Enrolment time Gestational age at entry (7-10 th gestation week); (30 Jun 2010 - 30 Jun 2011) Length Follow up until birth. Methods Clinic visits Data collection Questionnaire, medical records, laboratory measurements and diagnosis Loss to follow-up 0	second (21–24 gestational weeks) and third (33–37 gestational weeks) trimester of pregnancy. Every sample was assayed for TC, TG, HDL-C and LDL-C concentrations	Newborns were classified into appropriate for gestational age (AGA), SGA and LGA based on Neonatal Birth Weight for Gestational Age and Percentile in 15 Cities of China. LGA: birth weight above the 90 th percentile. SGA: birth weight below the 10 th percentile for gestational age. AGA: birthweight between LGA and SGA. According to the birth weight, neonates could be stratified into low birth weight (<2500 g), normal birth weight (2500–4000 g) and macrosomia (>4000 g) groups.	7
Tian et al. 2013	Study design: Prospective observational study	Setting: No statement Eligibility criteria: Maternal and neonatal characteristics were investigated between 2581 newborns with	No statement	No statement	Hypertriglyceridem ia and hypercholesterolem ia was diagnosed according to the	Macrosomia	Not applicab e

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
	Language: English Location: China	normal birth weight (controls,2500-3999g) and 306 macrosomia (birth weight over 4000g). Exclude criteria: Pregnancy with twins, premature labour and other complications were all excluded. Sample size: No statement			criteria of Hyperlipidaemia of National Cholesterol Education Program.		
Couch et al.1998	Study design: Perspective observational study Language: English Location: American	Setting: The Department of Obstetrics and Gynaecology, Hartford Hospital, Hartford, Connecticut, and private physicians' offices affiliated with Hartford Hospital Eligibility criteria: Women with GDM and healthy pregnant women with a negative diabetes screening test were recruited. Exclude criteria: Women with hypertension, hyperlipidaemia, renal or liver disease, heart disease, thyroid disorder, multiple gestations or parity >5 were excluded from the study. Sample size: n=40	Tx ± SD or n (%) Age (year) GDM: 31.6±2.7 Controls:30.6±3.2 Primiparous GDM: 8 (40%) Controls: 8 (40%) Maternal BMI (kg/m²) GDM:25.4±4.6 Controls:23.7±3.8 Gestational length GDM:38.3±1.7 Controls:37.6±2.2 Fasting blood Yes.	Enrolment time Gestational age at entry (26-30 th gestation week); (No statement on recruited time) Length Follow up until delivery Methods No statement Data collection Clinic diagnosis, clinic records. Loss to follow-up 0(0%)	Maternal plasma samples were collected between 37-38 gestation weeks and analysed for TC, HDL, LDL, VLDL and FFA	Cord vein samples were analysed for TC, HDL, LDL, VLDL and TG.	6
Ortega et al.1996	Study design: Cohort study Language: English Location: Spain	Setting: The INSALUD hospitals Eligibility criteria: Pregnant women carrying only a single child with no congenital malformations at 37 or more weeks of gestation. Participants without registered maternal disease (either before or during pregnancy), vaginal bleeding, blood pressure over 140/90 mm Hg, protein or glucose in the urine, pregnancy-related immunization and drug or alcohol abuse. Exclude criteria:	x ± SD or n (%) Age (year) 28.6±5.4 Primiparous NS Pre-pregnancy BMI (kg/m²) NS Gestational length 39.6±1.3 Fasting blood Yes.	Enrolment time Gestational age at entry (32-35 th gestation week); (October – December 1988) Length Follow up until delivery Methods Clinic visit Data collection Clinic diagnosis, obstetric case notes Loss to follow-up 0(0%)	Venous blood was collected at 32-35 gestation weeks after overnight fasting. TC, HDL-C, LDL-C, VLDL-C and triglycerides were measured by laboratory.	Birthweight was measured using a Marsden spring balance. Cord arteriovenous blood was obtained immediately after clamping and before delivery of the placenta. Blood samples were analysed for a series of lipid parameters (TC, HDL-C, LDL-C VLDL-C and triglycerides).	6

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		None. Sample size: n=292					
Alberti- Fidanza, et al.1995	Study design: Perspective observational study Language: English Location: Italy	Setting: Three towns in the Perugia area (Gubbio, Perugia and Umbertide) Eligibility criteria: Volunteer pregnant women attending the Maternity Advisory Service were recruited at the 1st trimester. Exclude criteria: Women and newborns in pathological conditions were not included. Sample size: n=70 For our interested association, the number of participants is 21.	No statement	Enrolment time Gestational age at entry (1st trimester); (No statement on recruited time) Length Follow up until 6 months post-partum Methods Clinic visits Data collection Laboratory measurements, clinic records, Loss to follow-up 49(70%)	At the 1 st , 2 nd and 3 rd trimester of pregnancy and at delivery, maternal venous blood was obtained for lipids assessments (TC, TG, HDL-C)	Mixed venous-arterial cord blood was obtained at delivery for TC, TG HDL-C measurements.	5
Brockerh off. 1986	Study design: Perspective observational study Language: Germany Location: German	Setting: Obstetrics Eligibility criteria: No statement Exclude criteria: No statement Sample size: n=112	No statement	No statement	Maternal blood was taken at 16 th gestation week for VLDL-C, LDL-C and HDL-C assessments.	Cord blood was obtained at delivery for TC and TG assessments.	
Robin et al. 2007	Study design: Retrospective cohort study Language: English Location: American	Setting: Hospital closest to the Greenwood Genetic Centre(GGC) in Greenwood, South Carolina Eligibility criteria: All women who were consecutively screened between 13 and 23 weeks' gestation during 1996-2001. Women who delivered at the hospital closest to GGC	x ± SD or n (%) Age (year) NS Primiparous NS Pre-pregnancy BMI (kg/m²) NS Gestational length	Enrolment time Gestational age at entry (No statement); (1996-2001) Length Follow up until delivery Methods Clinic visits Data collection	Maternal serum was taken between 13 and 23 weeks' gestation (mean:17.5 weeks, SD: 1.5 weeks) during 1996-2001. Frozen sera(-80°C)	Birthweight.	7

Study ID	Basic information	Participants	Maternal characteristics	Follow-up	Exposures	Outcomes	Quality score
		Exclude criteria: 1) Age<21 or >34 years old; 2) positive smoking history; 3) not dated by ultrasound 4) pregestational diabetes 5) twin gestation 6) race/ethnicity Hispanic, Asian, or Other 7) preeclamptic pregnancies 8) cardiac malformation 9) missing or conflicting data 10) foetal death 11) >1 eligible pregnancy to same mother 12) delivery before 37 gestation week Sample size: Low-TC group:100 Mid-TC group: 757 High-TC group:100	NS <u>Fasting blood</u> NS	Laboratory measurements, NIH clinical records, <u>Loss to follow-up</u> 47(9.9%) for low-TC group; 233(7.4%) for higher-TC group	were shipped on dry ice from GGC to the NIH. TC in serum was analysed in laboratory.		
Charles et al. 2016		Setting: Some centres (e.g. Malta) recruiting from a general population and others (eg. Greece and Italy) recruiting from an obstetric referral centre. Eligibility criteria: Pregnant Mediterranean women recruited in centres in Tunisia(n=112), Spain(n=187), Serbia(n=126), Malta(n=309), Italy(n=140), and Greece(n=178) who were not known to suffer from any form of carbohydrate metabolism problems outside their pregnancy (type 1 diabetes(T1DM), type 2 diabetes(T2DM), LADA, or MODY). Exclude criteria: None. Sample size: n=1062	$\frac{(kg/m^2)}{(m^2)}$	Enrolment time Gestational age at entry (27.9±2.3); (No statement on recruited time) Length Follow up until delivery Methods No statement Data collection Laboratory measurements, clinic records Loss to follow-up 0	Maternal fasting lipid profile levels were assayed at the time of the OGTT. Cholesterol, HDL-C, LDL-C and triglycerides were measured.	Birthweight.	5

S5 Appendix	Results	extract	ion forn
Study			

_	pendix Results exti	raction form					
Study				Maternal lipids			- Statistical Methods
ID		TC	HDL-C	LDL-C	TG	FFAs	
Ye et al.	$\overset{-}{x} \pm SD$ (mmol/L)	6.6 ± 1.4	2.4 ± 0.5	3.3 ± 0.8	2.9 ± 1.2	_	Statistical software: SPSS 16.0 Two tailed statistical tests and a significant p value < 0.05.
2015	Birth weight (g) (β, 95% CI)	9.1 (-6.4, 24.6)	-69.5 (-110, -28.2)	35.4 (10.1, 60.8)	25.2 (7.9, 42.6)	_	Multiple linear regression analysis adjusted for maternal glucose, maternal age, pre-pregnancy BMI, gestational weight gain, parity, neonatal sex and gestational age at delivery.
	SGA(n=39) (OR, 95% CI) AGA(n=873)	0.94 (0.74, 1.20) 1.00 (ref)	1.57 (0.87, 2.83) 1.00 (ref)	0.75 (0.50, 1.14) 1.00 (ref)	0.69 (0.47, 1.03) 1.00 (ref)	_	Logistic regression analysis adjusted for maternal age, pre- pregnancy BMI, gestational weight gain, parity and maternal
	LGA(n=331) (OR, 95% CI)	1.04 (0.94, 1.15)	0.62 (0.47, 0.82)	1.25 (1.06, 1.47)	1.15 (1.03, 1.27)	_	fasting blood glucose.
Wang et al.	Non-GDM (mmol/L) (Median, 25 th -75 th)	ND	$ \begin{array}{c} 1.88 \\ (1.65 - 2.12) \end{array} $	ND	1.95 (1.59 - 2.42)	_	Statistical software: SPSS 17.0
2015	GDM (mmol/L) (Median, 25 th -75 th)	ND	1.81 (1.50 – 2.09)	ND	$2.18 \\ (1.84 - 2.82)$	_	A significant p value < 0.05 .
	Birthweight (r, p)	ND	-0.12, p=0.01	ND	0.19, p<0.01	_	Partial correlation coefficients analysis adjusted for neonates' sex and gestational age.
Crume et al.	$(\bar{x} \pm SD, mg/dL)$	182.3±35.6	61.1±12.6	_	124.3±49.6	373.1±166.0	Statistical software: No statement
2015	$2^{\text{nd}} \underbrace{\text{visit}(20\text{-}34 \text{ week})}_{\text{(}}$ $\underbrace{\text{x} \pm \text{SD, mg/dL}}_{\text{)}}$	209.9±40.3	63.1±13.1	_	162.2±62.1	365.1±151.4	
	P value	< 0.0001	< 0.0001		< 0.0001	0.3	
	11-20 wk gestation Model 1 Birth	0.46±0.39 P=0.2	-0.54±1.17 P=0.6	_	0.09±0.30 P=0.7	0.06±0.09 P=0.5	Regression analyses were performed to determine the association of maternal metabolic fuels and metabolic
	weight $(\beta \pm SE, g,$ Model 2	0.42±0.42 P=0.3	-2.67±1.22 P=0.03	_	0.50±0.24 P=0.04	0.05±0.09 P=0.6	measures measured at each visit with neonatal outcomes. Model 1 adjusted for the residual value of the predictor from
	P) Model 3	0.44±0.41 P=0.3	-1.71±1.23 P=0.2	_	0.41±0.24 P=0.08	-0.11±0.10 P=0.2	the other visit, infant sex, gestational age at birth, maternal age, race/ethnicity, parity postnatal age at time of PEAPOD
	Birth Model 1	ND	-1.12±1.12 P=0.3	_	0.20±0.24 P=0.4	0.21±0.10 P=0.03	(for outcomes other than birth weight). Model 2 is model 1 plus maternal smoking, total energy intake, and maternal physical activity during pregnancy,
	$(x \pm SE, g, \beta, P)$ Model 2	ND	-3.12±1.16 P=0.07	_	0.39±0.24 P=0.1	0.31±0.11 P=0.003	gestational weight gain. Model 3 is model 2 plus pre-pregnancy BMI

Statistical Methods

Study				Maternal lipids	S		l
ID		TC	HDL-C	LDL-C	TG	FFAs	Ī
	Model 3	ND	-2.20±1.16 P=0.06	_	0.30 ± 0.24	0.24 ± 0.10	

The modification of effects of maternal cholesterol levels in late pregnancy on all neonatal body composition measures by pre-pregnancy BMI was reported in this study. A positive effect was noted for all neonatal outcomes (Birthweight, Fat mass, Fat free mass, Percent Fat mass) at higher pre-pregnancy BMIs, with a null effect for lean women and an inverse relationship on FM for underweight women. However, no β and P value around those associations was reported.

This study also reported that their findings were not influenced by the exclusion of women identified with GDM (n=26), gestational hypertension (n=61), or pre-eclampsia (n=34).

	GDW (n=20), gestatio	nai nypertension	(ii=01), or pre ceru	impsia (ii 5 i).			
Hwang et al.	$\frac{15-28 \text{ wks}}{(\text{x} \pm \text{SD, mg/dL})}$	_	_	_	143.4±68.5	_	Statistical software: SAS 9.3
2015	$\frac{29-42 \text{ wks}}{\text{x } \pm \text{SD, mg/dL}}$	_	_	_	273.4±123.3	_	Statistical significance was defined as P<0.05.
-	Birth weight (g), β (s.e.). p. R (%)					Maternal serum TG levels was log-transformed before
	15-28 wks	—	_	_	80.446 (31.738) P=0.0015, R=22.4	_	analyses due to its skewed distribution. Multiple regression analysis adjusted for maternal age, weight gain during
	29-42 wks	_	_	_	131.067 (31.242) P<0.0001, R=19.8	_	pregnancy, log-transformed urinary cotinine, gestational age, gestational age at blood collection, neonatal gender and long-transformed calorie intake.
Kulkar ni et	18 wks (x±SD,mmol/L)	4.11 ± 0.85	1.12 ± 0.28	_	1.09 ± 0.36	_	Statistical software: STATA version 11.2
al. 2013	28 wks (x±SD,mmol/L)	4.80 ± 0.89	4.80 ± 0.89	_	1.51 ± 0.52	_	
-	Birthweight (g): Mode	el 0 (\beta, 95% CI)					Model 0: Multiple regression analyses was performed to
	18 wks	39.07 (10.57, 67.58)	17.57 (-11.64, 46.77)	_	14.76 (-13.34 , 42.86)	_	explore the association of z-standardized maternal plasma glucose and lipid concentrations with neonatal measurements,
	28 wks	54.34 (24.85,83.88)	-8.89 (-38.72 ,20.95)	_	36.27 (4.32,68.23)	_	adjusting for gestation at the time of measurements, sex, SES, parity, maternal age, maternal BMI before pregnancy and total energy intake at the time of measurements.
	Birthweight (g): multi-	variate analyses	(β, 95% CI)				
	18 wks: model 1	33.42 (0.43,66.41)	6.68 (-24.08, 37.44)	_	4.24 (-26.40, 34.87)	_	Multiple analyses adjusted for gestation, sex of the baby, parity, SES, and maternal age, BMI before pregnancy, total
	28 wks: model 1	52.52 (19.11,85.92)	-21.58 (-52.62, 9.46)	_	23.93 (-11.29, 59.15)	_	energy intake at the time of measurements and other lipid levels.
	28 wks: model 2	44.42 (8.55,80.29)	-20.29 (-52.73, 12.14)	_	12.90 (-24.25, 50.06)	_	Model 1 entered with maternal fasting glucose. Model 2 entered with maternal 2-h glucose
Vrijkot	SGA (n=364)	4.97 ± 0.86	_	_	1.35 ± 0.61	_	Statistical software: SPSS 16.0 and the statistical package R

Study				Maternal lipids			— Statistical Methods
ID		TC	HDL-C	LDL-C	TG	FFAs	
te et al.	$(x \pm SD, mmol/L)$						2.13.1
2012	Non-SGA (n=3548) ($\bar{x} \pm SD$, mmol/L)	4.99 ± 0.87	_	_	1.33 ± 0.54	_	A P value <0.05 was considered statistically significant.
	$LGA (n=364)$ ($x \pm SD, mmol/L$)	5.06 ± 0.91	_	_	1.44 ± 0.61	_	
	Non-LGA (n=3548) ($\bar{x} \pm SD$, mmol/L)	4.98 ± 0.86	_	_	1.32 ± 0.54	_	
	Crude model						<u></u>
	SGA (OR, 95% CI)	0.97 (0.85-1.10)	_	_	1.06 (0.87-1.29)	_	Crude model: unadjusted associations between continuous TC and TG and the outcomes.
	LGA (OR, 95% CI)	1.10 (0.97-1.25)	_	_	1.44 (1.20-1.71)	_	and 10 and the outcomes.
	Model 1						Model 1 is multiple logistic regressions adjusted for maternal
	SGA (OR, 95% CI)	0.98 (0.86-1.12)	_	_	0.97 (0.79-1.19)	_	age, ethnicity, pre-pregnancy BMI, maternal education level, physical activity, smoking during pregnancy, and chronic
	LGA (OR, 95% CI)	1.08 (0.95-1.22)	_	_	1.48 (1.23-1.78)	_	hypertension.
Retnak aran et al. 2012	Lowest tertile birth weight infant [2020-3260 g] (n=156)	6.48 ± 1.25	1.73 ± 0.36	3.72 ± 1.17	2.25 ± 0.72	_	Statistical software: SAS 9.2
	Middle tertile birth weight infant [3260-3670 g] (n=157)	6.55 ± 1.23	1.72 ± 0.37	3.72 ± 1.12	2.46 ± 0.75	_	
	Highest tertile birth weight infant [3670-5700 g] (n=159)	6.39 ± 1.15	1.66 ± 0.34	3.6 ± 1.04	2.49 ± 0.66		
	p	0.5	0.2	0.5	0.006		Analysis of variance for continuous variables
	Birth weight (g, β,95 %	6 <u>CI)</u>					Multiple linear regression adjusted for length of gestation,
	Crude	ND	-120.54 (-244.42 to 3.35)	-15.22 (-55.49 to 25.05)	61.11 (-1.18 to 123.40)	_	infant sex, maternal demographic factors (age, ethnicity, family history of diabetes), smoking status, anthropometric measure (pre-pregnancy BMI, weight gain during pregnancy
	Adjusted	ND	-57.16 (-189.42 to 75.09)	-6.79 (-46.98 to 33.39)	-1.59 (-70.67 to 67.49)	_	up to the time of OGTT), glucose tolerance status, other lipid levels, insulin, adipokines (adiponectin, leptin) and inflammatory proteins (C-reactive protein)
	LGA (OR, 95% CI)						, r ,

Study				Maternal lipids	Ctatistical Watha Ja		
ID		TC	HDL-C	LDL-C	TG	FFAs	— Statistical Methods
	Crude	ND	0.89 (0.69 - 1.15)	0.80 (0.61 - 1.05)	1.26 (0.98 - 1.62)	_	Logistic regression analysis adjusted the same covariate as in the multiple linear regression analyses, except for length of
	Adjusted	ND	0.99 (0.70 - 1.39)	0.98 (0.72 - 1.34)	0.98 (0.70 - 1.38)	_	gestation and infant sex.
	White women LGA (O	R, 95% CI) (n=38					
	Crude	ND	0.82 (0.60 - 1.10)	0.85 (0.62 - 1.16)	1.33 (1.00 - 1.77)	_	
	Adjusted	ND	1.03 (0.69 - 1.52)	0.98 (0.69 - 1.38)	$ \begin{array}{c} 1.07 \\ (0.73 - 1.58) \end{array} $	_	Same statistical methods used in the LGA analyses.
Hou et al.	Mmol/L (median, 25 th -75 th)	6.28 (5.59-7.09)	1.75 (1.51-2.03)	3.06 (2.44-3.72)	3.05 (2.50-3.75)	_	Statistical software: SPSS 16.0 P<0.05 was considered statistically significant.
2014	AGA(n=2236)	6.30 (5.62-7.10)	1.76 (1.52-2.05)	3.07 (2.47-3.74)	3.02 (2.48-3.69)	_	Mann-whitney U test
	LGA(n=554)	6.18 (5.49-7.04)	1.70 (1.48-1.95)	2.95 (2.30-3.65)	3.19 (2.61-3.97)	_	
	p	0.017	0.000	0.003	0.000	_	
•	Outcome: LGA, (OR, 95% CI)						Binary logistic regression analyses adjusted for maternal age.
	Lowest tertile value	Ref	0.202 (0.026-1.562)	Ref	Ref	_	pre-pregnancy BMI, education level, smoking, annual household income, amniotic fluid volume, gestational
	Middle teritle value	0.967 (0.712-1.313)	Ref	0.785 (0.58-1.063)	3.037 (1.054-8.747)	_	hypertension, new-born sex, and gestational age at blood collection.
	Highest tertile value	1.084 (0.754-1.559)	0.812 (0.636-1.036)	0.829 (0.585-1.173)	3.303 (1.177-9.27)	_	The middle teritle value of maternal TC, HDL-C, LDL-C, TG and FFAs are 5.18-6.22, 1.04-1.55, 3.37-4.14 and 1.70-2.25.
	Infant weight gain at 3	3 months (β,p)					Statistical software: SAS 9.2
r et al. 2014	GDM group	-26.3,0.57	-150.6, 0.40	-11.7, 0.81	-43.3, 0.62	_	The unit of maternal lipid levels: mmol/L Multiple linear regression analyses adjusted for infant age at
	Non GDM group	37.0, 0.32	28.6, 0.80	43.5, 0.28	-14.2, 0.82	_	3-month visit, sex duration of exclusive breastfeeding, maternal and paternal ethnicity, birthweight and length of gestation.
Harmo	Mean \pm SEM				mg/dL	μEq/L	
n et al. 2011	Normal Early weight Late	_	_	_	85 ± 5.6	366 ± 52 326 ± 29	Statistical software: Sigama Stat for Windows version 2.03
	Obese Early Late	_	_	_	152 ± 14.3	535 ± 55 547 ± 58	
	None of the metabolic	measures correla	ted with birth we	ight (data not sho	wn).		A forward stepwise regression was used to generate models between infant adiposity and maternal metabolic parameters.

Study				Maternal lipid	s		Chatistical Wakes la
ID	_	TC	HDL-C	LDL-C	TG	FFAs	- Statistical Methods
Son et al.		$Mean \pm SD$	Mean ± SD		Median (IQR)		Statistical software: SPSS 12.0 (SPSS Inc., Chicago, IL USA)
2010	mmol/L	5.7 ± 1.1	1.7 ± 0.4	ND	2.5 (1.8-3.4)	_	p-value < 0.05 was considered significant.
	Non-LGA	5.8 ± 1.1	1.7 ± 0.5	ND	2.3 (1.8-3.1)		Differences between non-LGA group and LGA group wer
	LGA	5.5 ± 0.9	1.6 ± 0.3	ND	3.2 (2.4-3.6)		analysed using Student's t-test
	p	0.352	0.232	ND	0.001	_	_ analysed using Student 5 t test
	Birthweight (g, r, p)	p>0.05	p>0.05	ND	r = 0.17 p = 0.07	_	Statistical Method was not stated.
	LGA (OR, 95% CI)	ND	ND	ND	Hypertriglyceri demia (TG≥3.33 mmol/L) 4.43 (1.33-14.82)	_	Logistic regression model with confounding variables including parity, age, prepregnancy BMI, gestational weight gain.
Ahma d et al. 2006	Birthweight ratio (g, r ,p)	r = 0.147 p = 0.021	_	_	r = 0.122 p = 0.057	_	Birthweight ratio: birthweight adjusted for gestational age. Statistical software: SPSS 11.0. α =0.05, p<0.05 Univariate analysis.
					High TG (>2.78 mmol/L)		
	LGA (crude OR, 95% CI)	ND	_	_	3.07 (1.33, 7.08)	_	χ^2 test.
	LGA (adjusted OR, 95% CI)	ND	_	_	1.476 (1.15-1.93)	_	Backward wald mode in binary logistic regression. Adjuste for BMI, fasting plasma glucose and 2 hours postprandia plasma glucose.
Di et	$\text{mmol/L} (\bar{x} \pm \text{SD})$	6.34 ± 1.3	1.68 ± 0.4	4.01 ± 1	1.99 ± 0.64	_	Statistical software: SAS
al. 2005	birthweight (g, r ² , p)	ND	ND	ND	r ² =0.09 p<0.05	_	Univariate regression analyses.
					Hypertriglyceri demia (TG≥2.3 mmol/L)		χ^2 test.
	LGA (crude OR, 95%CI)	ND	ND	ND	5.6(0.93, 33.77)		
Schaef	_	mg/dL			mg/dL	μmol/L	Statistical software: SPSS 12.0 (Chicago, IL)
er- Graf et	$\frac{-}{x \pm SD}$ Week 28,32,36	253.7±55.6	_	_	265.9±87.6	262.6±112.4	All statistical tests were two-tailed and a P value <0.05 was considered significant.
al. 2008	Outcomes	ND	_	_	ND	ND	
2000	Close to delivery (r, p)						Bivariate correlation applying Spearman's correlation test

Study				Maternal lipid	S		- Statistical Methods
ID		TC	HDL-C	LDL-C	TG	FFAs	Stausucai Methods
	birthweight	ND	_	_	p>0.05	0.27, p=0.002	
	TGs in cord blood	ND	_	_	0.19, p=0.003	ND	
	FFAs in cord blood	ND	_	_	ND	0.28, p=0.004	
	After adjustment for m						
	from the profiles at 36				and TGs remained	independently	
	related to LGA (adjusted Maternal FFA levels w				nte than in mothers	with AGA	Logistic regression analysis
	infants (362.8 ± 101.7)			With LOA lines	nts than in mothers	with AGA	
	No statistically signific			parameters wit	h neonatal birth we	eight in the	Statistical software: PQStat software.
	GDM and NGT group			parameters with		and an and	P value <0.5 was considered statistically significant.
et al.							Multivariate linear regression for numerical factors and
2015							multivariate logistic regression were performed to assess the
Comm	- (SD)						influence of the factors affecting neonatal birth weight.
Somm er et	$mmol/L(x \pm SD)$ Visit 1	5.0 ± 0.9	1.73 ± 0.39	2.71 ± 0.73	1 21 + 0 55		Statistical software: IBM SPSS Statistics21, lincom
al.					1.31 ± 0.55	_	command in Stata IC 12
2015	Visit 2	6.2 ± 1.1	1.93 ± 0.45	3.44 ± 0.99	1.98 ± 0.69	_	
	Birthweight (g)						Data were provided by authors through email.
	Model 0 (β, 95%CI)	-4.2	-98.9	ND	48.8	_	W 110: : 1
	(р, услост)	(-39.4, 31.0)	(-188.1, -9.6)	1,2	(-14.8, 112.4)		Model 0 is simple regression analyses.
	Model 1 (β, 95%CI)	-6.1 (-37.5, 25.2)	-105.4 (-183.8, -27.0)	ND	94.4 (37.8, 150.9)	_	Model 1 is a multiple regression of the risk factor variables
		(-37.3, 23.2) -4.8	-118.8		(37.8, 130.9)		entered separately, adjusted for gestational week at inclusion.
	Model 2(β, 95%CI)	(-34.0, 24.4)	(-190.1, -47.5)	ND	(37.0, 133.7)	_	maternal age, parity, smoking status ethnic origin, offspring's
	N. 1.1.2 (0. 0.50 (CT)	-115.4	47.6) III	97.4		sex and gestational age.
	Model 3(β, 95%CI)	(-306.6, 75.8)	(-160.3, 255.6)	ND	(-3.8, 198.6)	_	Model 2 = Model 1 + early pregnancy BMI + weight gain.
	Model 4(β, 95%CI)	-74.9	-21.9	ND	83.4	_	Wodel 2 – Wodel 1 + early pregnancy BWI + weight gain.
	· ·	(-260.1, 110.2)	(-223.9, 180.2)	ND	(-14.6, 181.5)		Model 3: (risk variables are entered simultaneously into the
	Sum of skinfolds (mm)						regression, and adjusted for fasting glucose and 2-hour
	Model 0 (β, 95%CI)	0.17	-0.521	ND	0.583	_	glucose, maternal age, gestational week, parity, ethnicity
	V	(-0.14, 0.48) 0.10	(-1.312, 0.270) -0.608		(0.015, 1.151) 0.839		smoking status, offspring's sex and gestational age)
	Model 1 (β, 95%CI)	(-0.21,0.40)	(-1.381, 0.164)	ND	(0.280, 1.397)	_	Model 4 = Model 3 + early pregnancy BMI + weight gain.
	16 110/0 252/ 25	0.13	-0.611		0.724		inder i inder 5 i carry pregnancy Birit i weight gam.
	Model 2(β, 95%CI)	(-0.17,0.42)	(-1.321, 0.099)	ND	(0.245, 1.202)	_	
	Model 3(β, 95%CI)	-0.71	0.433	ND	0.623		
	Model 3(p, 9370Cl)	(-2.37, 0.95)	(-1.412, 2.279)	ND	(-0.308, 1.553)	_	

Study				Maternal lipids			- Statistical Methods
ID		TC	HDL-C	LDL-C	TG	FFAs	- Staustical Methods
	Model 4(β, 95%CI)	-0.44 (-2.08, 1.20)	-0.022 (-1.851, 1.808)	ND	0.577 (-0.341, 1.494)	_	
Slagja na et	$mmol/L (\bar{x} \pm SD)$ $LGA (n=50)$	6.0±1.0	1.3±0.4	3.8±1.0	3.8±1.8	_	Statistical software: SPSS 14.0 P<0.05 was considered statistically significant.
al.	AGA (n=135)	6.5±1.4	1.6 ± 0.4	3.5±1.2	3.1±1.1		, 0
2014	SGA (n=15)	6.3±1.3	1.5 ± 0.5	3.7±1.4	3.8±1.9		
	p (LGA vs. AGA)	p>0.05	0.001	p>0.05	0.012		Student t test
	p (AGA vs. SGA)	p>0.05	p>0.05	p>0.05	0.012		
	Birthweight (g, r, p)	ND	ND	ND	0.16, p=0.077	_	correlation analysis
	LGA (standardized β, p)	-0.230, p=0.164	ND	ND	0.326, p=0.045	_	Multiple linear regression
Laleh	$mg/dl (\bar{x} \pm SD)$						Statistical software: SPSS 16.0
t al.	28-32 wks	218.90±33.82	55.37±4.26	128.84 ± 29.23	175.71±24.23	_	
013	32-36 wks	240.99 ± 29.44	59.29±4.61	137.64 ± 29.22	240.46±32.06	_	
				1 15 10 00 50	252.05.20.61		
	A significant positive group after 32 weeks	of gestational age	(p<0.001) was for	und. (Bonferroni	multiple compariso	n test)	_
	A significant positive	correlation between gestational age ndependent predicts) was performed	een birth weight (L (p<0.001) was for action of birth weight. After adjustment	GA and macroso und. (Bonferroni ght in the study go for maternal pre	omia) and TG level is multiple comparison coup adjustment and	n test) alyses of	
Whyte	A significant positive group after 32 weeks of For determination of it covariance (ANCOVA only TG level remind	correlation between gestational age ndependent predicts) was performed	een birth weight (L (p<0.001) was for action of birth weight. After adjustment	GA and macroso und. (Bonferroni ght in the study go for maternal pre	omia) and TG level is multiple comparison coup adjustment and	n test) alyses of	Statistical software: SPSS 18.0
Whyte et al. 2013	A significant positive group after 32 weeks of For determination of it covariance (ANCOVA only TG level remind	correlation between gestational age ndependent predicts) was performed	een birth weight (L (p<0.001) was for action of birth weight. After adjustment	GA and macroso und. (Bonferroni ght in the study go for maternal pre	omia) and TG level is multiple comparison coup adjustment and	n test) alyses of	Statistical software: SPSS 18.0 A p value <0.05 was considered significant.
et al.	A significant positive group after 32 weeks of For determination of it covariance (ANCOVA only TG level remind mmol/L (x ± SD) Normal OGTT	correlation between of gestational age independent predicts. We was performed independently respectively.	een birth weight (Let (p<0.001) was for iction of birth weight. After adjustment lated to LGA (p=0)	GA and macroso und. (Bonferroni ght in the study go for maternal pre 1.04).	omia) and TG level in multiple compariso oup adjustment ana pregnancyBMI, ag	n test) alyses of	
et al.	A significant positive group after 32 weeks of For determination of it covariance (ANCOVA only TG level remind mmol/L (x ± SD) Normal OGTT (n=167) Abnormal OGTT	correlation between of gestational age ndependent predicts. A) was performed independently researched by the control of the co	een birth weight (Le (p<0.001) was for action of birth weight. After adjustment lated to LGA (p=0)	GA and macrosound. (Bonferroni ght in the study grant for maternal presented).	omia) and TG level is multiple comparison coup adjustment ana -pregnancyBMI, ago	n test) alyses of	
et al.	A significant positive group after 32 weeks of For determination of it covariance (ANCOVA only TG level remind mmol/L (x ± SD) Normal OGTT (n=167) Abnormal OGTT (n=22)	correlation between of gestational age ndependent predicts. A) was performed independently researched by the control of the co	een birth weight (Le (p<0.001) was for action of birth weight. After adjustment lated to LGA (p=0)	GA and macrosound. (Bonferroni ght in the study grant for maternal presented).	omia) and TG level is multiple comparison to adjustment anal-pregnancyBMI, agrants 1.84±0.86	n test) alyses of	
et al.	A significant positive group after 32 weeks of For determination of it covariance (ANCOVA only TG level remind mmol/L (x ± SD) Normal OGTT (n=167) Abnormal OGTT (n=22) Birthweight (kg) (x ±	correlation between of gestational age ndependent predicts. A) was performed independently researched by the control of the co	een birth weight (Le (p<0.001) was for action of birth weight. After adjustment lated to LGA (p=0)	GA and macrosound. (Bonferroni ght in the study grant for maternal presented).	omia) and TG level is multiple comparison coup adjustment ana pregnancyBMI, age 1.84±0.86 2.33±0.78 mmol/L	n test) alyses of	
et al.	A significant positive group after 32 weeks of For determination of it covariance (ANCOVA only TG level remind mmol/L (x ± SD) Normal OGTT (n=167) Abnormal OGTT (n=22) Birthweight (kg) (x ± <2.99 3.09-3.49 3.5-3.99	correlation between of gestational age ndependent predicts. A) was performed independently researched by the control of the co	een birth weight (Le (p<0.001) was for action of birth weight. After adjustment lated to LGA (p=0)	GA and macrosound. (Bonferroni ght in the study grant for maternal presented).	nmia) and TG level is multiple comparison to adjustment analy-pregnancyBMI, age 1.84±0.86 2.33±0.78 mmol/L 1.58±0.40 1.88±0.93 1.87±0.73	n test) alyses of	
et al.	A significant positive group after 32 weeks of For determination of it covariance (ANCOVA only TG level remind mmol/L (x ± SD) Normal OGTT (n=167) Abnormal OGTT (n=22) Birthweight (kg) (x ± <2.99 3.09-3.49 3.5-3.99 4.0-4.49	correlation between of gestational age ndependent predicts of a second predicts of the control o	een birth weight (Le (p<0.001) was for iction of birth weight. After adjustment lated to LGA (p=0) 1.54±0.41 1.39±0.35	.GA and macroso und. (Bonferroni ght in the study gu for maternal pre 1.04). 2.74±0.78 2.86±0.75	nmia) and TG level is multiple comparison to adjustment anarpregnancyBMI, age 1.84±0.86 2.33±0.78 mmol/L 1.58±0.40 1.88±0.93 1.87±0.73 2.23±1.119	n test) allyses of e, and parity, — — — — — — — — — — — — — — — — — —	A p value <0.05 was considered significant.
et al.	A significant positive group after 32 weeks of For determination of it covariance (ANCOVA only TG level remind mmol/L (x ± SD) Normal OGTT (n=167) Abnormal OGTT (n=22) Birthweight (kg) (x ± <2.99 3.09-3.49 3.5-3.99 4.0-4.49 Maternal triglyceride	correlation between of gestational age ndependent predicts of a performed independently residue of the second of t	een birth weight (Le (p<0.001) was for iction of birth weight. After adjustment lated to LGA (p=0) 1.54±0.41 1.39±0.35	.GA and macroso und. (Bonferroni ght in the study grater for maternal present). 2.74±0.78 2.86±0.75	mia) and TG level is multiple comparison to up adjustment analopregnancyBMI, age 1.84±0.86 2.33±0.78 mmol/L 1.58±0.40 1.88±0.93 1.87±0.73 2.23±1.119 crease in birth weight	n test) alyses of e, and parity, — — — — — — ht (p<0.03).	A p value <0.05 was considered significant. Univariate analysis
et al.	A significant positive group after 32 weeks of For determination of it covariance (ANCOVA only TG level remind mmol/L (x ± SD) Normal OGTT (n=167) Abnormal OGTT (n=22) Birthweight (kg) (x ± <2.99 3.09-3.49 3.5-3.99 4.0-4.49	correlation between of gestational age ndependent predicts. A) was performed independently response to the second	een birth weight (Le (p<0.001) was for iction of birth weight. After adjustment lated to LGA (p=0) 1.54±0.41 1.39±0.35 ———————————————————————————————————	.GA and macroso und. (Bonferroni ght in the study grater for maternal present). 2.74±0.78 2.86±0.75	mia) and TG level is multiple comparison to up adjustment analopregnancyBMI, age 1.84±0.86 2.33±0.78 mmol/L 1.58±0.40 1.88±0.93 1.87±0.73 2.23±1.119 crease in birth weighting increased birthweighting multiple series and series are series and series are series and series are series and series are	n test) alyses of e, and parity, — — — — — — th (p<0.03). ght (p<0.04).	A p value <0.05 was considered significant. Univariate analysis
et al.	A significant positive group after 32 weeks of For determination of it covariance (ANCOVA only TG level remind mmol/L (x ± SD) Normal OGTT (n=167) Abnormal OGTT (n=22) Birthweight (kg) (x ± <2.99 3.09-3.49 3.5-3.99 4.0-4.49 Maternal triglyceride of Maternal increased tri	correlation between of gestational age ndependent predicts. A) was performed independently response to the second	een birth weight (Le (p<0.001) was for iction of birth weight. After adjustment lated to LGA (p=0) 1.54±0.41 1.39±0.35 ———————————————————————————————————	.GA and macroso und. (Bonferroni ght in the study grater for maternal present). 2.74±0.78 2.86±0.75	mia) and TG level is multiple comparison to up adjustment analopregnancyBMI, age 1.84±0.86 2.33±0.78 mmol/L 1.58±0.40 1.88±0.93 1.87±0.73 2.23±1.119 crease in birth weighting increased birthweighting multiple series and series are series and series are series and series are series and series are	n test) alyses of e, and parity, — — — — — — th (p<0.03). ght (p<0.04).	A p value <0.05 was considered significant. Univariate analysis Multivariate regression analysis adjusting for age, BMI and

Study				Maternal lipids			- Statistical Methods
ID		TC	HDL-C	LDL-C	TG	FFAs	- Statistical Methods
2012							difference between groups.
	Normal BW (n=890)	6.05 ± 1.53	2.20 ± 0.45	2.76 ± 0.71	2.43 ± 1.48	_	
	p	>0.05	< 0.05	>0.05	>0.05	_	
	Hypo-HDL-cholestero	<u>lemia</u>					Unconditional logistic regression model.
	Crude OR	ND	1.67	ND	ND	_	
	Adjusted OR (95%CI)	ND	1.63(1.02-2.60)	ND	ND	_	Adjusted for maternal age and BMI.
	p <u>Macrosomia</u>	ND	0.04	ND	ND	_	
	HDL-c (mmol/L) >2.49	Case (all, %) 14 (234, 6.0%)	OR (95%CI)	p 			HDL-C was categorized in quartiles based on the distribution in all pregnant women, and risk in each quartile was
	2.18-2.49		1.59(0.78-3.27)	0.202			estimated in reference to lowest or highest quartile of
	1.87-2.16		1.47(0.72-2.99)	0.291			metabolic marker level.
	<1.87		2.09(1.04-4.21)	0.039			
Vrijkot	$\text{mmol/L} (\bar{x} \pm \text{SD})$, ,	,				Statistical software: SPSS 16.0
te et	Birth weight<2500g	4.63±0.79		_ (1.21±0.56	_	A P value<0.05 was considered significant.
al.	2500g-4000g	4.97±0.86		_	1.31±0.53	_	Ç
2011	Birth weight>4000g	5.01±0.89	_	_	1.40±0.62	_	
•	Standardised Birthwei	$ght, \beta(SE)$					Standardized birthweight (already adjusted for gestational age
	TC(mmol/L)	Univariate	Model 1	TG (mmol/L)	Univariate	Model 1	at birth, parity and sex)
	Q1 (3.87±0.33)	-0.12 ± 0.07	-0.09 ± 0.06	Q1	-0.03±0.07	-0.06±0.06	Univariate associations between TG and TC levels and BW
	$Q2(4.48\pm0.13)$	0.07 ± 0.07	0.09 ± 0.06	Q2	0.03 ± 0.07	0.00 ± 0.06	SDS were explored by using regression analyses.
	$Q3(4.89\pm0.12)$	Reference	Reference	Q3	Reference	Reference	Model 1 is multivariate analyses further adjusted for maternal age, maternal height, hypertension, maternal pre-pregnancy
	$Q4(5.36\pm0.15)$	0.07 ± 0.07	0.08 ± 0.06	Q4	0.04 ± 0.07	0.03±0.06	BMI, weight gain during early pregnancy, ethnicity, smoking,
	Q5(6.23±0.61)	0.11 ± 0.07	0.11 ± 0.06	Q5	0.17 ± 0.07	0.20 ± 0.06	alcohol use, education level, and cohabitant status
•	Standardised Birthwei	ght,					Data were provided by authors through email.
	β(95%CI)	11.82 (-10.00, 33.65)	_	_	47.14 (12.42, 81.87)	_	Univariate linear analysis
	β(95%CI)	22.67 (4.00, 41.33)	_	_	86.72 (56.13,117.30)	_	Multivariate results linear analysis adjusted for maternal age, maternal height, hypertension, maternal pre-pregnancy BMI, weight gain during early pregnancy, ethnicity, smoking, alcohol use, education level, and cohabitant status
•	SDS weight						Linear regression analyses were used to exploring
	A significantly differe infants born of women the growth patterns of	with the lowest	TG levels (Q1) de	viated more from	their individual	growth line than	associations between different TG and TC quintiles and postnatal growth patterns (weight, length, and BMI expressed as SDS).

Study				Maternal lipids			- Statistical Methods
ID		TC	HDL-C	LDL-C	TG	FFAs	Staustical Methods
	progressively increase analyses showed that differences were 0.140 SDS length The individual average there was a tendency for differences at 1 months. SDS BMI A similar tendency was 12 months. Differences Accelerated weight gas. The percentage of infa was significantly high length and BMI shows between Q1 and other. No associations were and no post hoc differences weight for gestational Differences between TD Differences between TD Differences between TD Differences between TD Differences were 1.140 SDS Length for gestational Differences between TD Differences Descriptions and DIFFERENCE DIFFERE	differences in wei O SDS for Q1 vs Q e lines with SDS for the Q1 pattern i only for Q1 vs Q as observed for Q es were present at tin ants in Q1 that sho er compared with ed a similar tende TG quintiles wer found between TG ences). I age according to TG quintiles: %SQ	ght among TG qu Q5, and 0.139 SDS did not differ sign to deviate (p=0.00 Q5 (0.140). 1, with a relatively the first month after the other TG quin ncy with regard to re found. C quintiles and we TG and TC quint GA (p=0.768), %L	intiles were only in the for Q1 vs Q3. If for Q1 vs Q3. If ficantly among s 61). Post hoc analy low BMI at 1 m ter birth only for growth (24.5%) do an accelerated go ight, length and Hiles: GA (p=0.032)	ubsequent TG qui lyses revealed sign onth and a relative Q1 vs Q3 (0.129). uring the first 6 m %; P=0.027). Althrowth, no significa	onth; these Intiles, although hificant Ely high BMI at the south of life hough both ant differences	A multivariable model adjusted for maternal age, maternal height, parity, maternal pre-pregnancy BMI, weight gain during early pregnancy, ethnicity, education level, cohabitant status, smoking, alcohol use, pregnancy duration, infants' age and BW. To compare SDS trajectories between the TG and TC quintiles in more detail, post hoc comparisons were done at multiple time points: 1, 3, 6, 9 and 12 months. The amount of accelerated growth in the different quintiles was determined by using the Pearson χ^2 analysis.
Vinod	Gestational age-adjus	ted birth weight (g) - Normal weigh	<u> it group – β(95%</u>	<u>CI)</u>		Statistical software: SAS 9.1
et al.			mg/	/dL			A p value of <0.05 was considered significant.
2011	6-10 wks (n=62)		-4.1 (-10.4, 2,2)		1.1 (-0.4, 2.6)	_	Univariate regression analyses.
	10-14 wks (n=65)		-2.1 (-7.7, 3.6)		1.5 (0.1, 2.8)	_	
	16-20 wks (n=68)		-1.0 (-6.4, 4.4)	` ' '	0.7 (-0.8, 2.1)	_	
	22-26 wks (n=71)		-4.1 (-8.8, 0.6)	` ' '	1.1 (0.0,2.1)	_	
	32-36 wks (n=69)		-3.6 (-8.6, 1.4)		0.9 (-0.1, 1.9)	_	
	Gestational age-adjus	-	-				
	6-10 wks (n=69)		-7.7 (-16.1, 0.7)		0.4 (-2.3, 3.0)	_	
	10-14 wks (n=71) 16-20 wks (n=65)	1.5 (-1.8, 4.7)	-8.0 (-15.6, -0.4) -9.3 (-16.4, -2.1)	1 1	1.4 (-0.5, 3.2) 0.7 (-1.2. 2.6)	_	
	22-26 wks (n=71)		-9.5 (-16.4, -2.1) -7.4 (-14.1,-0.7)		1.5 (0.1, 3.0)		
	32-36 wks (n=70)		-10.0(-17.5, -2.3)		1.9 (0.6, 3.2)		
	The effect size of mate			1 1			
	HDL quartile	$\frac{-}{x\pm}$ SD (mg/dL)	Mean differen	~			
	Normal weight	-		_			

Study				Maternal lipid	s		- Statistical Methods
ID		TC	HDL-C	LDL-C	TG	FFAs	- Stausucai Methods
	1(lowest)	60.3±3.5	Refer	ence			
	2	70.4 ± 3.0	-36.5 (-86				
	3	80.5 ± 2.8	72.7 (-173	3.7, 28.3)			
	4	100.3±11.5	-144 (-3	44,56)			
	Obese/Overweight						
	1(lowest)	60.0±4.1	Refer				
	2	68.8±1.9	-88 (-154				
	3	79.1±4.3	-191 (-334				
	4	94.7±8.2	-347 (607.	.3, -79.8)			
Zawiej	mmol/L	_	1.87(1.59,2.26)	_	2.45(3.22,4.24)	_	Statistical software: SPSS 12.0
ska et	(Median, 25 th -75 th)						P<0.05 was considered statistically significant.
al. 2008	Birthweight (g)		ND		$R^2 = 0.02$ F = 9.43		Linear regression analyses.
2000	R^2 , F, p		ND	· 7	P = 9.43 P < 0.01	_	Linear regression analyses.
					1 < 0.01		Data were provided by the author through email.
	Macrosomia		0.59(0.32,1.02)				Population: non-obese GDM women
	(RR,95%CI, p)		P=0.051		ND		T
							Chi-square statistics.
Clause	mmol/L	5.3(4.8,5.9)	1.8 (1.5,2.0)	2.8 (2.3,3.3)	1.5 (1.2,1.9)		Statistical software: SPSS 11.0
n et al.	(median, 25 th -75 th)		1.0 (1.3,2.0)	2.0 (2.3,3.3)	1.3 (1.2,1.9)		P<0.05 was considered statistically significant.
2005	Macrosomia (OR, 95%						
	Triglycerides (case/all)	v	Model A	Model B	Model C	Model D	
	Q1 (10/437)	1.0	1.0	1.0	1.0	1.0	
	Q2 (28/668)	1.9 (0.9-3.9)	1.7(0.8-3.6)	1.9(0.9-3.9)	1.6(0.7-3.3)	1.4(0.7-3.1)	Q, quartile
	Q3 (15/394)	1.7(0.8-3.8)	1.4(0.6-3.2)	1.7(0.7-3.8)	1.4(0.6-3.2)	1.3(0.5-2.9)	Univariate logistic regression was used to calculate
	Q4 (35/551)	2.9(1.4-5.9)	2.2(1.1-4.6)	2.9(1.4-5.9)	2.5(1.2-5.2)	1.9(0.9-4.1)	unadjusted OR value.
	P trend	0.004	0.062	0.004	0.016	0.121	Multiple logistic regression analyses was performed in Mode
	TC (case/all)	unadjusted OR	Model A	Model B	Model C	Model D	A, B, C and D. Variables in model A: first trimester BMI;
	Q1 (20/497)	1.0	1.0	1.0	1.0	1.0	Model B: age, parity smoking
	Q2 (19/565)	0.8(0.4-1.6)	0.8(0.4-1.5)	0.8(0.4-1.6)	0.7(0.4-1.4)	0.7(0.3-1.3)	Model C: age, parity smoking, weight gain, placental weight,
	Q3 (25/448)	1.4(0.8-2.6)	1.4(0.7-2.5)	1.4(0.8-2.5)	1.3(0.7-2.4) 0.9(0.5-1.7)	1.4(0.7-2.6) 0.9(0.5-1.7)	gestational diabetes
	Q4 (24/540) P trend	1.1(0.6-2.0) 0.397	1.0(0.5-1.8) 0.610	1.1(0.6-2.0) 0.451	0.9(0.5-1.7)	0.9(0.5-1.7)	Model D: model C+ first trimester BMI
	, i	*					
	HDL-C(case/all) Q1 (38/509)	unadjusted OR 1.0	Model A 1.0	Model B 1.0	Model C 1.0	Model D 1.0	

Study				Maternal lipids			- Statistical Methods
ID		TC	HDL-C	LDL-C	TG	FFAs	- Statistical Methods
	Q2 (18/498)	0.5(0.3-0.8)	0.5(0.3-0.9)	0.5(0.3-0.8)	0.3(0.3-0.9)	0.6(0.3-1.0)	
	Q3 (18/527)	0.4(0.2-0.8)	0.5(0.3-1.0)	0.4(0.2-0.7)	0.5(0.2 - 0.8)	0.3(0.3-1.0)	
	Q4 (14/516)	0.3(0.2-0.6)	0.4(0.2-0.8)	0.3(0.2-0.6)	0.4(0.2-0.7)	0.4(0.2-0.8)	
	P trend	< 0.001	0.008	< 0.001	0.001	0.009	
	Non-HLD-C(case/all)	unadjusted OR	Model A	Model B	Model C	Model D	
	Q1 (16/519)	1.0	1.0	1.0	1.0	1.0	
	Q2 (19/530)	1.2(0.6-2.3)	1.2(0.6-2.3)	1.2(0.6-2.3)	1.0(0.5-2.0)	1.0(0.5-2.1)	
	Q3 (21/500)	1.4(0.7-2.7)	1.3(0.7-2.5)	1.4(0.7-2.7)	1.2(0.6-2.5)	1.3(0.7-2.7)	
	Q4 (32/499)	2.2(1.2-4.0)	1.9(1.0-3.5)	2.1(1.2-3.9)	1.8(1.0-3.5)	1.9(1.0-3.6)	
	P trend	0.009	0.034	0.011	0.036	0.035	
Mathe	mmol/L (median, 5th -	9 th)					Statistical software: SPSS 10.0
ws et al.	Early pregnancy (n=733)	5.59(4.30,7.45)	_		_	_	P<0.05 was considered statistically significant. P value cautiously throughout and considered value <0.05 but >0.0
2003	Later pregnancy (n=537)	6.91(5.30,9.14)	_	- 0	_	_	as marginal
	Birthweight (g, β, 95%	6CI <u>)</u>			NL.		_
	Early pregnancy (≈16wks, n=733)	30.1(1.21.58,9) P=0.041	_	_	\-\P	_	Multiple linear regression model adjusted for matern smoking status and height, infant' gender, gestational age.
	Later pregnancy (≈28wks n=537)	11.1(-18.0, 40.3) P= 0.453	_	_	- 16	917.	shoking status and neight, infant gender, gestational age.
Olmos	$\text{mmol/L}(\bar{x} \pm \text{SD})$						Statistical software: PASW statistics version 18.00, GraphPa
et al. 2014	2 nd trimester _Normal weight	ND	ND	_	1.99±0.65	_	Prism 5.0 for Windows. P<0.05 was considered statistically significant.
	2 nd trimester _Overweight	ND	ND	_	2.29±0.75	_	
	2 nd trimester _Obese	ND	ND	_	2.35 ± 0.71	_	
	3 rd trimester _ Normal weight	ND	ND	_	2.59±0.76	_	
	3 rd trimester _ Overweight	ND	ND	_	2.76±0.91	_	
	3 rd trimester _ Obese	ND	ND	_	2.88 ± 0.92	_	
	Newborn weight z-sco	re (r, p)					Maternal lipids z score – newborn weight z score
	Normal weight (n=128)	ND	ND	_	r=0.12,p=0.158	_	Linear regression model.
	Overweight (n=105)	ND	ND	_	r=0.42,p<0.001	_	
	Obese (n=46)	ND	ND	_	r=0.47,p<0.001	_	

Study				Maternal lipids			— Statistical Methods
ID		TC	HDL-C	LDL-C	TG	FFAs	— Statistical Methods
Emet	$mg/dL(\bar{x} \pm SD)$						Statistical software: SPSS 15.05
et al.	1 st trimester	166.20 ± 28.28	53.37±10.51	93.75 ± 23.22	93.09±45.57	_	P<0.05 was considered statistically significant.
2013	3 rd trimester	271.28±47.81	63.54±21.16	154.58±44.15	274.10±101.89	_	_
	Birthweight (p)	0.616	0.754	0.440	0.033	_	Changed maternal lipid levels - birthweight
	Neonatal weight in 3 rd postnatal month (p)	0.2678	0.860	0.769	0.138	_	Pearson correlation analyses.
Liu et al.	$\text{mmol/L}(\bar{x} \pm \text{SD})$						Statistical software: SPSS 17.00 P<0.05 was considered statistically significant.
2016	GDM	6.09 ± 0.86	1.82 ± 0.35	3.26 ± 0.86	2.31±0.84	_	
	NGT	3.30 ± 0.81	1.85 ± 0.33	3.30 ± 0.81	2.09 ± 0.76	_	
	Birth weight (r, p)	0.018, p=0.518	-0.011, p=0.701	-0.005, p=0.843	0.100, p<0.001	_	Partial correlation adjusted for gestational age and pre-gravid BMI
	Birthweight (β , SE, p)	ND	ND	ND	0.070, SE=13.235 P=0.001	_	Multiple linear regression model including First Visit FPG, OGTT FPG, triglyceride, Apolipoprotein E, pre-gravid BMI, GDM, gestational age.
Brunn er et al.	$mg/dL (\bar{x} \pm SD)$	_	_	_	197.0±66.2	_	Statistical software: R version 2.8.1, PASW version 18.0. A tow-sided P-value<0.05 was considered statistically significant.
2013	Maternal lipid levels a	t gestation weeks	32 (β,95%CI)				
	Birthweight(g)	_	_	_	-0.54 (-1.56, 0.49)	1/-;	Data were provided by authors through email.
	Ponderal index (kg/m³)	_	_	_	-0.00 (-0.01, 0)		Multiple linear regression model, including the covariates
	6 weeks postpartum weight (g)	_	_	_	-0.97 (-2.33, 0.4)	_	maternal pre-pregnancy BMI, gestational weight gain, maternal glucose tolerance status, pregnancy duration, sex
	6 weeks postpartum ponderal index (kg/m³)	_	_	_	-0.00 (0, 0)	_	and group allocation for the data at birth, and, additionally, poderal index at birth and mode of infant feeding at the later
	4 months postpartum weight (g)	_	_	_	-0.62 (-2.27, 1.03)	_	time points, were performed.
	4 months postpartum ponderal index (kg/m³)	_	_	_	0.01 (0, 0.01)	_	
	1 year postpartum weight (g)	_	_	_	-1.46 (-3.83, 0.92)	_	
	1 year postpartum ponderal index (kg/m³)	_	_	_	-0.00 (-0.01, 0)	_	
	1 year postpartum	_	_		-0.00	_	

Study	_			- Statistical Methods			
ID		TC	HDL-C	LDL-C	TG	FFAs	— Statistical Methods
	BMI (kg/m ²)				(0, 0)		
	No significant relations	nips were found	l for maternal trigl	lyceride levels a	t 32 nd gestation week	with	
	birthweight and Ponders	al index (or BM	I) at delivery, 6 w	eeks, 1 years an	nd 2 years post-partui	n, and also	
	with weight gain after b	irth at any time	point.	•			
	The change in maternal	serum triglycer	ide concentration	between the 15th	th and 32 nd week of go	estation was	
	weakly, but significantly)
						0.001 (0-0.01)
			41	1 1 : 4:	:		
		ot with any of th	ne other growth or	body compositi	ion outcomes up to 2	years post-	
	partum.	ot with any of th	ne other growth or	body compositi	ion outcomes up to 2	years post-	
Cnopp		ot with any of th	ne other growth or	body compositi	ion outcomes up to 2	years post-	Statistical software: No statement.
et al.	partum.	ot with any of th	ne other growth or	body compositi	ion outcomes up to 2	years post-	Statistical software: No statement.
	partum. $mM (\bar{x} \pm SD)$	t with any of th	ee other growth or	body compositi	•	years post-	Statistical software: No statement.
et al.	partum. $mM (\bar{x} \pm SD)$ NS-(n=521)	t with any of th	ee other growth or — — — —	body compositi	1.86±0.68	years post-	Statistical software: No statement.
et al.	partum. $mM (\bar{x} \pm SD)$ NS- (n=521) PS+ (n=264)	— — — — — — — — — — — — — — — — — — —	e other growth or — — — —	body compositi	1.86±0.68 1.92±0.68	years post-	Statistical software: No statement. Univariate Spearman's correlation coefficients
et al.	partum. $mM (\bar{x} \pm SD)$ NS- (n=521) PS+ (n=264) GDM (n=96)	— — — — — — — — — — — — — — — — — — —	ee other growth or — — — — —	body compositi	1.86±0.68 1.92±0.68	years post-	
et al.	partum. mM (x ± SD) NS- (n=521) PS+ (n=264) GDM (n=96) <u>Birthweight ratio</u>	— — — — — — — — — — — — — — — — — — —	ee other growth or — — — — — — —	body composition	1.86±0.68 1.92±0.68 2.29±0.68	years post-	
et al.	partum. mM (x ± SD) NS- (n=521) PS+ (n=264) GDM (n=96) Birthweight ratio NS-	— — — — — — — — — — — — — — — — — — —	ee other growth or — — — — — — — — — — — — — — —	body composition	1.86±0.68 1.92±0.68 2.29±0.68 0.09 (p≤0.05)	years post-	
et al.	partum. mM (x ± SD) NS- (n=521) PS+ (n=264) GDM (n=96) Birthweight ratio NS- PS+	— — — — — — — — — — — — — — — — — — —	ee other growth or	body composition	1.86±0.68 1.92±0.68 2.29±0.68 0.09 (p≤0.05) 0.13(p≤0.05)	years post-	

	Birthweight ratio						Univariate Spearman's correlation coefficients
	NS-	_	_	_	$0.09 (p \le 0.05)$	_	
	PS+	_	_	_	$0.13(p \le 0.05)$	_	
	GDM	_	_	_	0.11	_	
	PS+ plus GDM	_	_	_	$0.16(p \le 0.01)$	_	
	ALL	_	_	_	$0.12(p \le 0.01)$	_	
Knopp			HDL-C	LDL-C	VLDL-C	FFAs	
et al.	Separman pairwise cor	relation coeffici	<u>ents</u>				Spearman rank correlation coefficients indicate the linear
1985	Birth weight (n=273)	_	-0.06	0.003	0.05	-0.06	relationship between all pairs of variable.
	Birth weight ratio (n=248)	_	-0.06	0.01	0.03	0.002	
	Standardized regressio	n coefficients					Structured multiple regression analyses. Variables in very unit were entered the regression equation sequentially and in
	Birth weight (n-272)	_	-0.15	0.04	-0.14	0.05	a predefined order. Unit I: VLDL-C, VLDL-TG,LDL-C, HDL-TG
	Birth weight ratio (n=247)	_	-0.13	0.01	-0.30, p<0.05	-0.09	Unit II: Glucose, insulin, FFA, HPL, progresterone, estradiol and estriol
Schaef		mmol/L			mmol/L	μmol/L	Statistical software: SPSS 16.0
er-	$\bar{x} \pm SD$	6.56 ± 0.11	_	_	2.84 ± 0.08	320±14	P<0.05 was considered statistically significant.
Graf et al. 2011	A significant lineal post transformed FFAs (r=0 None of the maternal n	.1886, p=0.0172	<i>)</i> .			or log	Pearson correlation coefficients were calculated to evaluate the correlations between different variables.
Nolan	TG(r, p)	Asian-born	GDM(n=38)	Asian & GDM	Overall		Statistical software: SPSS-PC software package

Study				Maternal lipids			- Statistical Methods
ID		TC	HDL-C	LDL-C	TG	FFAs	- Staustical Methods
et al.		(n=97)		(n=18)	(n=388)		
1995	Birth weight ratio (univariate analyses)	0.23, p=0.02	0.37, p=0.023	0.63, p=0.005	0.12, p=0.02	_	All statistical tests were two-tailed, and a P value of <0.05 was considered significant.
	Birth weight ratio (multiple regression)	ND	P=0.004	ND	ND	_	Within the total GDM subgroup, using multiple regression analyses to control for the maternal factors of BMI and rate of maternal weight gain.
Friis et al.	$mmol/L(\bar{x} \pm SD)$	6.96±1.20	1.71±0.37	_	2.01±0.65	0.44±0.13	Statistical software: SPSS 18.0. All p-values ,0.05 were considered statistically significant.
2012	Birthweight (β,95%CI, p)	p>0.05	-170 (-329, -9) P=0.04	_	94(2,187) P=0.046	p>0.05	Multiple linear regression model adjusted for gestational age at birth.
Lei et	mmol/L(median, IQR)	_	1.46 (1.3-1.7)	_	2.71(2.12-3.49)	_	Statistical software: SPSS 22.0.
al. 2016	OR (95%CI)	TG (<3.49 mmol/L)	TG (≥3.49 mmol/L)	<i>HDL-C</i> (≥1.3 mmol/L)	<i>HDL-C</i> (<1.3 mmol/L)		Logistic regression.
	LGA	1 (Ref)	1.6 (1.42-2.01)	1 (Ref)	1.33(1.12-1.58)	_	
	SGA	1 (Ref)	1.51(1.08-2.12)	1 (Ref)	0.88(0.62-1.25)		
Kitaji ma et al.		mg/dL			mg/dL	mEq/dL	Statistical software: SAS 5.0
2001	$\bar{x} \pm SD$	263.6±46.2	_	_	213.9±77.7	70.3±12.3	P<0.05 was defined as significant
	Birthweight (r, p)	0.01, p=0.99	_	_	0.22, p=0.009	0.03, p=0.73	Univariable linear regression.
	Birthweight (F,p)	ND	_	_	6.3, p=0.014	ND	After controlling for fasting plasma glucose, prepregnant BMI, maternal weight gain during pregnancy, gestational age at delivery, neonatal gender.
		Hypertriglyceri demia	Normal triglyceride	p	Crude OR	?(95%CI)	χ^2 test
	LGA Non-LGA	4 30	1 111	0.012	14.8 (1.59	, 137.38)	
	LGA Hypertriglyceridemia	Adjusted OR 11.6	95%CI (1.1 - 122)	p 0.04			Logistic regression model adjusted for fasting plasma glucose levels, prepregnant BMI, and weight gain during pregnancy
Mossa	$mg/dL (\bar{x} \pm SD)$	201.4±38.4	46.6±4.36	115.3±34.9	197.5±51.9	_	Statistical software: SPSS 20.0
yebi et	Birthweight (g)						P<0.05 was defined as significant
al. 2014	r, p	0.50, p<0.001	-0.47, p<0.001	0.40, p<0.001	0.68, p<0.001	_	Pearson correlation analyses.
2014	β, SE	ND	ND	ND	5.24, SE=0.54	_	Stepwise linear regression adjusted for male gender of the child
	Standardized β , p	ND	ND	ND	0.59, p<0.001	_	

Study				Maternal lipids			Statistical Mathada
ID	•	TC	HDL-C	LDL-C	TG	FFAs	Statistical Methods
	<u>Macrosomia</u>				TG	TG z score	Forward stepwise logistic regression analyses
	β, SE, p	ND	ND	ND	0.04, SE=0.01 P<0.001	ND	Adjusted for maternal age, weight prior to pregnancy, FE and cholesterol.
	OR (95% CI)	ND	ND	ND	1.044(1.02-1.07)	9.44(2.86-31.16)	and cholesterol.
	<u>LGA</u>						Forward stepwise logistic regression analyses
	β, SE, p	ND	ND	ND	0.03, SE=0.01 P<0.001	ND	Adjusted for maternal age, weight prior to pregnancy, Fl
	OR (95% CI)	ND	ND	ND	1.035(1.02, 1.05)	5.90 (2.68-13.00)	and cholesterol.
	<u>LGA</u>	<u>all</u>	Case(proportion)	Crude OR(95%CI)	aOR (95%CI)		
	Total cholesterol:						Logistic regression model
	Q1:<172	39	2 (5.1)	1 (Ref)	1 (Ref)		
	Q2:172.1-199.9	35	6 (17.1)	3.8 (0.7-20.4)	2.3 (0.4-15.2)		
	Q3:200-234.9	37	9 (24.3)	5.9 (1.2-29.7)	1.2 (0.2-8.6)		
	Q4:≥235	43	18 (41.9)	13.3 (2.8-62.5)	1.1 (0.2-8.1)		
	HDL:						
	Q1: ≤43	40	18 (45.0)	16.4 (3.5-77.2)	0.6 (0.07-5.3)		
	Q2:43.1-46	37	10 (27.0)	7.4 (1.5-36.5)	0.08 (0.08-5.6)		
	Q3:46.1-49.9	35	5 (14.3)	3.3 (0.6-18.4)	1.7 (0.2-11.6)		Variables in model: mother's age, weight prior to pregnan
	Q4: ≥50	42	2 (4.8)	1 (Ref)	1 (Ref)		FBS, triglyceride, cholesterol, and child gender. If
	<u>LDL:</u>						categorical variable was one of these confounders or l
	Q1: <88	38	3 (7.9)	1 (Ref)	1 (Ref)		colinearity with other variables, we excluded that varia
	Q2:88.1-113	40	9 (22.5)	3.4 (0.8-13.6)	2.04 (0.4-10.9)		and only the categorical variable was entered.
	Q3:113.1-143.9	37	10 (27)	4.3 (1.1- 17.3)	0.6 (0.1-4.03)		
	Q4: ≥144	39	13 (33.3)	5.8 (1.5-22.6)	0.8 (0.1-4.4)		
	<u>Triglyceride:</u>						
	Q1: <170	37	2 (5.4)	1 (Ref)	1 (Ref)		
	Q2:170-199.9	37	0 (0)	0	0		
	Q3:200-299.9	37	6 (16.2)	3.4 (0.6-18)	3.2 (0.5-20.7)		
	Q4: ≥230	43	27 (62.8)	29.5 (6.2-139.6)	28.2 (3.5-230.3)		
_	mmol/L (median, IQR))					Statistical software: SPSS 20.0
al.	Early pregnancy (n=284)	4.58 (3.87-5.39)	0.64(0.46-0.97)	3.31(2.66-3.94)	1.31(0.80-1.35)	_	
2016	Late pregnancy (n=293)	6.02(5.00-6.87)	0.85(0.54-1.13)	4.15(3.43-5.06)	1.71(1.28-2.19)	_	
	Early pregnancy						×:p>0.1, statistically insignificant;

Study			Maternal lipids			Challed and Made also
ID	TC	HDL-C	LDL-C	TG	FFAs	— Statistical Methods
Birth weight	×	×	×	×	_	√:p<0.1, statistically significant
Sum of skinfold	×	×	×	$\sqrt{}$	_	Pearson correlation was used, and Spearman's correlation for
2 year weight centile	$\sqrt{}$	×	×	×	_	the nonparametric data to individually measure the
2 years old waist: length ratio	×	$\sqrt{}$	×	×	_	correlation between each blood lipid (in early and late pregnancy and cord blood), HOMA, C-peptide and leptin
2 years old sum of skinfold	×	×	$\sqrt{}$	×	_	concentration and each of the anthropometric measures of child weight and adiposity (at birth, 6 months and 2 years of age).
<u>Late pregnancy</u>						Bivariate associations at a significance of $P < 0.1$ were
Birth weight	×	×	×	$\sqrt{}$	_	considered significant
Sum of skinfold	×	$\sqrt{}$	×	$\sqrt{}$	_	constacted significant
2 year weight centile	$\sqrt{}$	×	$\sqrt{}$	$\sqrt{}$	_	
2 years old waist: length ratio	×	\checkmark	×	×	_	
2 years old sum of skinfold	×	×	×	×	_	
Birthweight (g) (β, p, 95%CI)	ND	ND	ND	β=111.17 p=0.034 (8.48, 213.87)	_	Multiple regression model controlling for confounders (at birth: mother's BMI, gestational age, infant gender, mother's education and smoking status, and at 6-month and 2-years:
Birthweight centile	×	×	×	(0.10, 213.07) √	_	infant gender, age at data collection, mother's education
2 years old weight	×	×	×	×	_	status and breastfeeding), outcomes associated with maternal
Subgroup analyses_ la						blood parameters were birth weight, birth weight centile, and
Birthweight (BMI< 25kg/m²)	ND	ND	ND	R ² =0.0003, p=0.92	_	weight at 6 months. The final multiple linear regression models that were
Birthweight (BMI≥25kg/m²)	ND	ND	ND	R ² =0.08, P=0.008	_	statistically significant ($P < 0.05$) were reported as the best predictors of infant weight and adiposity.
Birthweight(g) (β,95%	6CI <u>)</u>					Data were provided by authors through email.
Early pregnancy	27.87 (-17.89,73.63)	-1236.25 (-3322.95, 850.45)	18.39 (-38.44, 75.21)	ND	_	Multiple regression model (controlling for mother's BMI, gestational age, infant gender, mother's education and
Late pregnancy	24.85 (-9.39, 59.09)	30.00 (-114.85, 174.84)	19.97 (-24.34, 64.27)	111.18 (8.48, 213.87)	_	smoking status)
<u>Sum of skinfolds (β,95</u>						
Early pregnancy	0.23 (-0.96, 1.41)	-1.59 (-5.68, 2.51)	0.19 (-1.19, 1.56)	ND	_	
Late pregnancy	0.61 (-0.49, 1.71)	-0.16 (-4.24, 3.92)	0.46 (-0.74, 1.66)	ND	_	
Weight at 2 years(kg)	<u>(β,95%CI)</u>					Multiple regression model (controlling for infant gender, age

Study				Maternal lipids			- Statistical Methods
ID		TC	HDL-C	LDL-C	TG	FFAs	- Staustical Methods
	Early pregnancy	0.15	0.24	0.12	0.71		at data collection, mother's education status and
	Early pregnancy	(-0.14, 0.44)	(-0.82, 1.29)	(-0.23, 0.47)	(-0.06, 1.48)		breastfeeding)
	Late pregnancy	0.23	0.16	0.27	0.47		
Τ' ,		(-0.02, 0.48)	(-0.77, 1.09)	(-0.05, 0.58)	(-0.05, 0.99)		and the appearance
Jin et al.	mmol/L (median, IQR	*	1 (((1 45 1 77)	2.25 (2.09.2.45)	2 20 (1 77 2 72)		Statistical software: SPSS 19.0
2016	1 st (7-10 weeks) 2 nd (21-24 weeks)	3.95 (3.66-4.60) 4.65 (4.22-5.10)	1.66 (1.45-1.77) 1.67 (1.47-1.79)	2.25 (2.08-2.45)	2.20 (1.77-2.73) 2.45 (2.11-2.89)	_	P values < 0.05 were defined as statistically significant.
2010	3 rd (33-37 weeks)	4.63 (4.22-3.10) 6.27 (5.52-7.03)	1.80 (1.57-2.04)	2.46 (2.22-2.77) 2.87 (2.32-3.45)	3.06 (2.37-3.98)	_	
	1st trimester (Adjusted		1.80 (1.57-2.04)	2.87 (2.32-3.43)	3.00 (2.37-3.98)		_
	` 3	, , , , , ,	1.31 (0.32-5.38)				
	SGA	ND	P=0.709	ND	ND	_	Forward stepwise logistic regression analysis.
	Macrosomia	ND	0.51 (0.19-1.36) P=0.178	ND	ND	_	Odds ratios were adjusted for maternal age, prepregnancy BMI, gestational weight gain, parity, maternal education
	2 nd trimester (Adjusted	d OR, 95%CI, p)					background, family income and cigarette exposure. Values of
	SGA	ND	1.88 (0.47-7.59) P=0.377	ND	ND		macrosomia and SGA were additionally corrected for delivery mode and infant sex.
	Macrosomia	ND	0.25 (0.09-0.73) P=0.011	ND	ND	_	
	3 rd trimester (Adjusted						
	SGA	1.12 (0.80-1.56) P=0.520	3.15 (1.15-8.65) P=0.026	1.16 (0.71-1.89) P=0.565	0.63 (0.40-0.99) P=0.046	7.	Odds ratios were adjusted for maternal age, prepregnancy BMI, gestational weight gain, parity, maternal education
	LGA	0.98 (0.86-1.11) P=0.715	0.79 (0.52-1.21) P=0.281	0.93 (0.78-1.11) P=0.418	1.13 (1.02-1.26) P=0.025	40	background, family income and cigarette exposure. Values of PTB, SGA, LGA and macrosomia were additionally
	Macrosomia	0.99 (0.81-1.21) P=0.903	0.46 (0.22-0.94) P=0.034	0.93 (0.69-1.25) P=0.621	1.19 (1.02-1.39) P=0.024	_	corrected for delivery mode and infant sex.
Tian et	OR (95%CI)				\geq 2.27mmol/L		No statement on statistic software and method.
al. 2013	Macrosomia	_	_	_	2.20 (1.54-3.14)	_	
Couch et al. 1998	In control group, mate In control group, mate significantly correlate In GDM group, mater	ernal HDL-C sign d with cord vein I	ificantly correlate FFAs (r=0.47, p≤0	d with cord vein 70.05).	$\Gamma C (r=0.51, p \le 0.05)$. Maternal TG	Software: Statistical Analysis Systems Program Pearson correlation analyses
Ortega	U 1,	TC	HDL-C	LDL-C	TG	VLDL-C	Statistical software: No statement
et al. 1996	mmol/L ($\bar{x} \pm SD$) Newborn lipids (r, p)	6.82±1.16	1.62±0.34	4.07±1.07	2.43±0.83	1.11±0.38	P<0.05 were considered to indicate statistical significance.
	TC HDL-C	0.3298, p<0.05 0.2575, p<0.05	ND ND	0.3204, p<0.05 ND	ND ND	ND ND	Spearman's rank correlation
		5.20 / C, P (0.00		.,.	.,2		

Study			Maternal lipids			— Statistical Methods
ID	TC	HDL-C	LDL-C	TG	FFAs	— Statistical Methods
LDL-C	0.3053, p<0.05	ND	0.3507, p<0.05	ND	ND	
TG	ND	ND	ND	ND	ND	
VLDL-C	ND	ND	ND	ND	ND	
mmol/L	Maternal TC<7	.55 mmol/L	Maternal TC≥7.	55 mmol/L	n	Student t test.
	(n=21	5)	(n=77)	p	Student t test.
$TC(\bar{x} \pm SD)$	1.65±0	.47	2.10±0.	54	< 0.05	
$HDL-C(\bar{x} \pm SD)$	0.63±0	0.25	0.75±0.	21	< 0.05	
LDL-C($\bar{x} \pm SD$)	0.78 ± 0	0.36	1.14±0.	40	< 0.05	
$TG(\bar{x} \pm SD)$	0.48 ± 0	0.22	$0.45\pm0.$	20	>0.05	
$VLDL-C(\bar{x} \pm SD)$	0.22±0	0.10	0.21±0.	09	>0.05	
TC/HDL-C($\bar{x} \pm SD$)	2.62±0	.40	2.81±0.	35	< 0.05	
Birthweight (g, $\bar{x} \pm SD$)	3301.5±	406.6	3234.5±4	11.5	>0.05	

Alberti Maternal HDL-C levels in the 2nd trimester is significantly associated with cord blood triglycerides among all nowborns (r=-0.53, p=0.0131, n=21).

Fidanz For girls (n=7), maternal HDL-C levels in the 1st (r=-0.86, p=0.0138) and 2nd (r=-0.83, p=0.0218) trimester a, et al. is significant associated with cord blood TG respectively. Maternal triglycerides measured in the 2nd trimester is correlated with cord blood TC level (r=0.80, p=0.0315). No correlation was observed among boys.

Pearson linear correlation.

Brocke rhoff	r, p Cord blood TC		HDL 0.484	LDL 0.082	VLDL 0.828, P<0.01		No statement on statistic methods.
1986	Cord blood TG		0.063	0.246	0.568, P<0.01		
Robin et al. 2007	Birthweight, g	Mean(SD)	Unadjusted mean difference, p	Adjusted mean difference, p			Unadjusted mean difference was assessed using 1-way analysis of variance, comparing low-TC or high-TC group with mid-TC reference group. Adjusted mean difference was assessed using multivariate
	Mid-TC group	3484(482)	Ref	Ref			linear regression; model adjusted for infant gender, fractional
	Low-TC group	3360(442)	-124, 0.015	-150, 0.001			week of GA within the term interval, maternal weight in pounds, maternal age group, and race in pooled analyses.
	High-TC group	3504(471)	+20, 0.69	+29, 0.47			Outliers measurement were excluded from the adjusted model.
Charle s et al. 2016	Birthweight (r, p)	-0.103, p<0.0001	-0.139, p<0.0001	0.001, p<0.0001	-0.014, p<0.0001	_	No statement on statistical software as well as statistical significant level. Pearson correlation.

ND: No documented.

S6 Appendix Quality assessment form

S6 Appendix Quality a	assessn				G	7 474				
Study ID		Selec		A 4		rability		Outcom		Overall Score
Harmon et al.2011	A1 0	A2	A3	A4	B1 0	B2 0	C1 0	C2	C3	5
Son et al.2010	0	1	1	1	0	0	0	1	1	5
Di et al.2005	0	1	1	1	0	0	0	1	1	5
Schaefer-Graf et al.2008	0	1	1	1	0	0	0	1	1	5
Slagjana et al.2014	0	1	1	1	0	0	0	1	1	5
Zhou et al.2012	0	1	1	1	0	0	0	1	1	5
Zawiejska et al.2008	0	1	1	1	0	0	0	1	1	5
Emet et al.2013	0	1	1	1	0	0	0	1	1	5
Schaefer-Graf et al.2011	0	1	1	1	0	0	0	1	1	5
Mossayebi et al.2014	0	1	1	1	0	0	0	1	1	5
Swierzewska et al.2015	0	1	1	1	0	0	0	1	1	5
Ortega et al.1996	0	1	1	1	0	0	0	1	1	5
Alberti-Fidanza et al.1995	0	1	1	1	0	0	1	1	0	5
Charles et al. 2016	0	1	1	1	0	0	0	1	1	5
Wang et al.2015	0	1	1	1	1	0	0	1	1	6
Ahmad et al. 2006	0	1	1	1	1	0	0	1	1	6
Whyte et al. 2013	0	1	1	1	0	0	1	1	1	6
Vinod et al. 2011	0	1	1	1	1	0	0	1	1	6
Olmos et al.2014	0	1	1	1	1	0	0	1	1	6
Knopp et al.1992	0	1	1	1	1	0	0	1	1	6
Nolan et al.1995	0	1	1	1	1	0	0	1	1	6
Friis et al.2012	0	1	1	1	1	0	0	1	1	6
Lei et al.2016	0	1	1	1	1	0	0	1	1	6
Kitajima et al.2001	0	1	1	1	1	0	0	1	1	6
Couch et al.1998	0	1	1	1	0	0	1	1	1	6
Brockerhoff 1986 Retnakaran et al.2012	0	1	1	1	0	0	$\frac{1}{0}$	1	1	6 7
Hou et al.2014	0	1	1	1	1	1	0	1	1	7
Laleh et al.2013	0	1	1	1	1	1	0	1	1	7
Liu et al.2016	0	1	1	1	1	0	1	1	1	7
Brunner et al.2013	0	1	1	1	1	0	1	1	1	7
Knopp et al.1985	0	1	1	1	1	0	1	1	1	7
Geraghty et al.2016	0	1	1	1	1	1	0	1	1	7
Jin et al.2016	0	1	1	1	1	1	0	1	1	7
Robin et al. 2007	0	1	1	1	1	1	0	1	1	7
Ye et al.2015	0	1	1	1	1	1	1	1	1	8
Crume et al.2015	0	1	1	1	1	1	1	1	1	8
Hwang et al.2015	0	1	1	1	1	1	1	1	1	8
Kulkarni et al.2013	1	1	1	1	0	1	1	1	1	8
Vrijkotte et al.2012	0	1	1	1	1	1	1	1	1	8
Kramer et al.2014	0	1	1	1	1	1	1	1	1	8
Vrijkotte et al. 2011	0	1	1	1	1	1	1	1	1	8
Clausen et al.2005	1	1	1	1	0	1	1	1	1	8
Mathews et al.2003	0	1	1	1	1	1	1	1	1	8
Sommer et al.2015	1	1	1	1	1	1	1	1	1	9

S7 Appendix Data analysis for birthweight

Data summary

S7.1 Table Results summary of the association of maternal lipid levels with birthweight throughout pregnancy

Maternal lipids	Trimester	Negative associations	No direction	Positive associations	Total
	The first trimester	1	1	2(1)	4
TC	The second trimester	1	4	7(2)	12
	The third trimester	3(1)	12	8(3)	23
	The first trimester	2(1)	0	0	2
HDL-C	The second trimester	6(2)	4	1	11
	The third trimester	11(6)	6	1	18
	The first trimester	1	0	1	2
LDL-C	The second trimester	1	5	2	8
	The third trimester	2	5	7(3)	15
	The first trimester	0	1	4(3)	5
TG	The second trimester	0	2	10(8)	12
	The third trimester	3(1)	4	20(14)	27
	The first trimester	0	0	0	0
VLDL	The second trimester	0	0	0	0
	The third trimester	0	1	1	2
	The first trimester	0	1	0	1
FFAs	The second trimester	0	0	1	1
	The third trimester	0	3	4(2)	7

^{1.} This table summarised the results distribution of studies that reported the association of maternal lipid levels with birthweight throughout pregnancy;

^{2.} Number in this table represent the number of studies;

^{3. &#}x27;No direction' means that the number of studies reported statistically insignificant results without its direction, as well as the number of studies did not report their results;

^{4.} Number in the bracket means the number of studies reported statistically significant results;

^{5.} Abbreviation: Total cholesterol (TC), high-density lipoprotein cholesterol (HDL-C), low-density lipoprotein cholesterol (LDL-C), very low-density lipoprotein cholesterol (VLDL-C), triglycerides (TG) and free fatty acids (FFAs).

Total cholesterol (TC)

ID	Population	Countries	Sample Tri.	Reported		Lower		n	Statistical	Quality	The	cont	rol of	f conf	ound	ling	fact	ors
ID.	1 opulation	Countries	size	measures		95%CI	95%CI	p	methods	Quanty	a	b	c	d	e	f	g	h
Vinod et al.2011(1)	Normal weight	USA	65 1	Crude β	-19.33	-120.03	81.36	ND	SLR	6	×	×	×	×	×	×	×	×
Vinod et al.2011(2)	Overweight/obese	USA	71 1	Crude β	58.00	-67.86	183.87	ND	SLR	6	×	×	×	×	×	×	×	×
Vrijkotte et al.2011	General	Netherlands	2,052 1	Crude β	11.82	-10.00	33.65	ND	Univariate analyses	8			×	×	×	×	√	×
Vrijkotte et al.2011	General	Netherlands	2,052 1	Adjusted β	22.67	4.00	41.33	ND	MLR	8		$\sqrt{}$	$\sqrt{}$			×		×
Nolan et al.1995	General	Australia	388 1	ND	ND			ND	ND	6	ND	ND	ND	ND	ND	ND	×	ND
Liu et al.2016	General	China	1,546 2	r	0.02			0.518	Partial correlation	7	×	×	×	×	×	×	×	×
Vinod et al.2011(1)	Normal weight	USA	71 2	Crude β	-50.27	-112.24	11.69	ND	SLR	6	×	×	×	×	×	×	×	×
Vinod et al.2011(2)	Overweight/obese	USA	71 2	Crude β	3.87	-91.02	98.75	ND	SLR	6	×	×	×	×	×	×	×	×
Mathews et al.2003	General	UK	733 2	Adjusted β	30.10	1.21	58.90	ND	MLR	8	$\sqrt{}$	$\sqrt{}$	×	×	×	×	$\sqrt{}$	×
Crume et al.2015	General	USA	804 2	Adjusted β	17.79	-11.82	47.39	0.200	MLR	8	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	×	×	×	$\sqrt{}$	×
Kulkarni et al.2013	non-GDM	India	631 2	Adjusted β	39.07	10.57	67.58	ND	MLR	8	×	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	×	×	$\sqrt{}$	×
Geraghty et al.2016	non-GDM	UK	331 2	Adjusted β	27.87	-17.89	73.63	ND	MLR	7	$\sqrt{}$	$\sqrt{}$	×	$\sqrt{}$	$\sqrt{}$	×	×	×
Whyte et al. 2013	General	Ireland	189 2	ND	ND			ND	ND	6	ND	ND	ND	ND	ND	ND	×	ND
Wang et al.2015	General	China	636 2	ND	ND			ND	Partial correlation	6	$\sqrt{}$	$\sqrt{}$	×	×	×	×	×	×
Di et al.2005	OGTT+	Italy	83 2	ND	ND			ND	ND	5	ND	ND	ND	ND	ND	ND	×	ND
Olmos et al.2014	GDM	Chile	279 2	ND	ND			ND	ND	6	ND	ND	ND	ND	ND	ND	×	ND
Mossayebi et al.2014	General	Iran	154 3	r	0.50			<0.001	Pearson correlation	5	×	×	×	×	×	×	$\sqrt{}$	×
Charles et al. 2016	General	Multiple	1062 3	r	-0.103			<0.0001	Pearson correlation	4	×	×	×	×	×	×	×	×
Ahmad et al. 2006	non-GDM	Malaysia	246 3	r	0.16			0.021	Univariate analyses	6	$\sqrt{}$	×	×	×	×	×	$\sqrt{}$	×
Kitajima et al.2001	OGTT +	Japan	146 3	r	0.01			0.990	SLR	6	×	×	×	×	×	×	$\sqrt{}$	×
Vinod et al.2011(1)	Normal weight	USA	69 3	Crude β	-46.40	-118.05	25.24	ND	SLR	6	×	×	×	×	×	×	×	×
Vinod et al.2011(2)	Overweight/obese	USA	70 3	Crude β	15.47	-89.10	120.03	ND	SLR	6	×	×	×	×	×	×	×	×
Sommer et al.2015	General	Norway	699 3	Crude β	-4.20	-39.40	31.00	ND	SLR	9	×	×	×	×	×	×		×
Sommer et al.2015	General	Norway	699 3	Adjusted β	-6.10	-37.50	25.20	ND	MLR	9	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	×	×	×		×
Mathews et al.2003	General	UK	537 3	Adjusted β	11.10	-18.00	40.30	ND	MLR	8			×	×	×	×		×
Ye et al.2015	non-GDM	China	1,243 3	Adjusted β	9.10	-6.40	24.60	ND	MLR	8		$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	×

ID	Population	Countries	Sample Tri.	Reported	Effect Lower		p	Statistical	Quality	The	cont	trol of	f conf	found	ling	fact	
		Countries	Size	measures	size 95%C			methods		a	b	c	d	e	f	g	h
Kulkarni et al.2013	non-GDM	India	631 3	Adjusted β	54.34 24.8	85 83.88	ND	MLR	8	×	√	√	V	×	×	V	×
Geraghty et al.2016	non-GDM	UK	331 3	Adjusted β	24.85 -9.3	59.09	ND	MLR	7	$\sqrt{}$	$\sqrt{}$	×	$\sqrt{}$		×	×	×
Couch et al.1998	General	USA	40 3	p	ND		>0.05	Pearson correlation	6	×	×	×	×	×	×	×	×
Ortega et al.1996	General	Spain	292 3	p	ND		>0.05	Student t test	5	×	×	×	×	×	×	$\sqrt{}$	×
Swierzewska et al.2015	General	Poland	136 3	p	ND		>0.05	MLR	5	ND	ND	ND	ND	ND	ND	×	ND
Emet et al.2013	General	Turkey	801 3	p	ND		0.616	Pearson correlation	5	×	×	×	×	×	×	×	×
Friis et al.2012	General	German	207 3	p	ND		>0.05	MLR	6	$\sqrt{}$	×	×	×	×	×	×	×
Retnakaran et al.2012	non-GDM	Canada	472 3	p			0.500	Analysis of variance for continuous variables	7	×	×	×	×	×	×	×	×
Schaefer-Graf et al.2011	non-GDM	German	190 3	p	ND		>0.05	Pearson correlation	5	×	×	×	×	×	×	$\sqrt{}$	×
Son et al.2010	GDM	Korea	104 3	p	ND		>0.05	ND	5	ND	ND	ND	ND	ND	ND	$\sqrt{}$	ND
Crume et al.2015	General	USA	804 3	ND	ND		ND	MLR	8	$\sqrt{}$	$\sqrt{}$	\checkmark	×	×	×	$\sqrt{}$	×
Slagjana et al.2014	non-GDM	Yugoslavia	200 3	ND	ND		ND	ND	5	ND	ND	ND	ND	ND	ND	×	ND
Olmos et al.2014	GDM	Chile	279 3	ND	ND		ND	ND	6	ND	ND	ND	ND	ND	ND	×	ND
Schaefer-Graf et al.2008	GDM	German	150 3	ND	ND		ND	Spearman correlation	5	×	×	×	×	×	×	×	×
Robin et al. 2007	General	American	957 2		Adjusted MD(g)	MLR	7	$\sqrt{}$		\checkmark	×	×	×		×
			High-TC gro	oup (n=100)	Ref group	Ref g	group										
			Mid-TC gro	oup(n=757)	29	0.	47										
			Low-TC gro	oup(n=100)	-150	0.0	001										

The bold font represents statistically significant results.

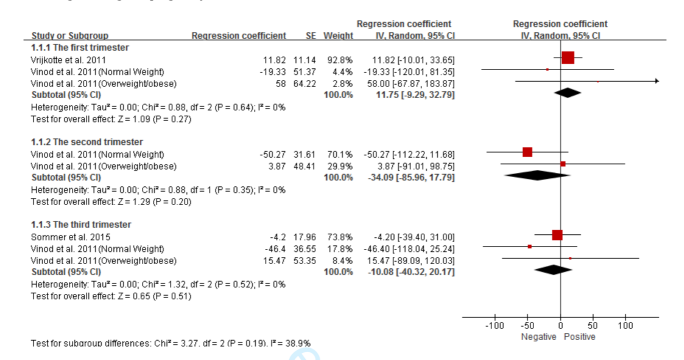
Confounding factors: a. Gestational age; b. Neonatal gender; c. Maternal age; d. Pre-pregnancy BMI; e. Gestational weight gain; f. Maternal glucose level; g. pre-term birth; h. Maternal lipid levels.

Abbreviation: Trimesters(Tri.), Gestational diabetes mellitus(GDM), Positive screenes of Oral Glucose Tolerance test(OGTT+), Confidence interval(CI), Not documented(ND), Simple linear regression(SLR), Multiple linear regression(MLR), United Kingdom(UK), Mean difference(MD), Reference(Ref).

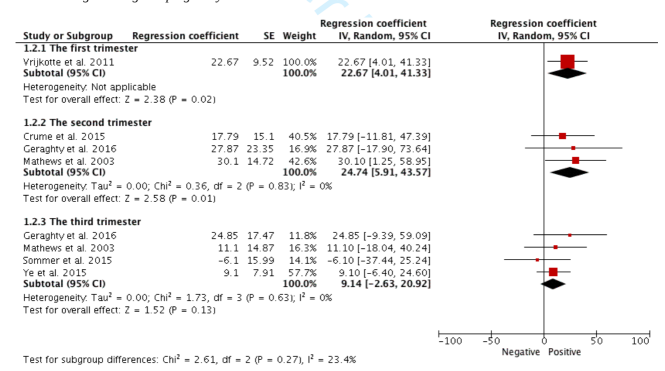
r: Correlation coefficients; β: regression coefficients.

Meta-analysis

S7.1 Figure Overall meta-analysis of crude regression coefficients for the association between maternal TC levels and birthweight throughout pregnancy

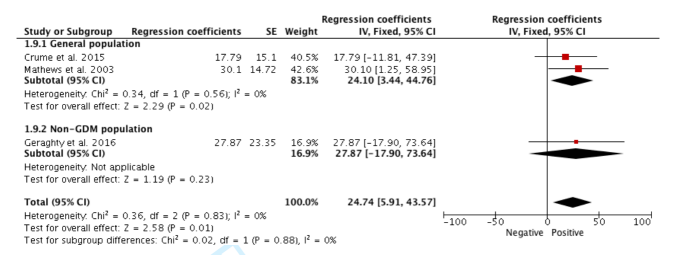


S7.2 Figure Overall meta-analysis of adjusted regression coefficients for the association between maternal TC levels and birthweight throughout pregnancy

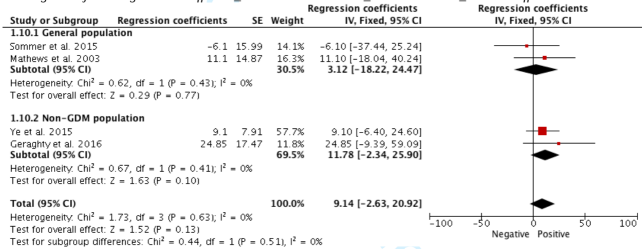


Subgroup analysis

S7.3 Figure Adjusted regression coefficient_ General vs. non-GDM_ the 2nd trimester_ Random effect model

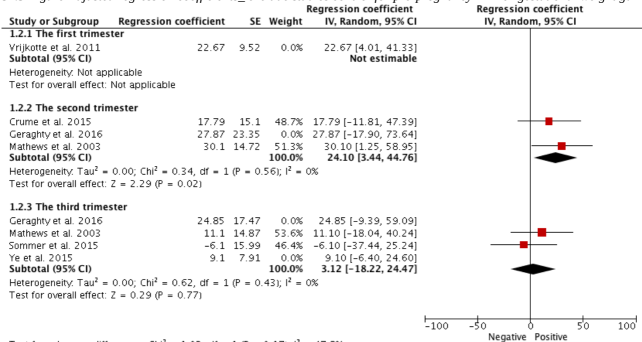


S7.4 Figure Adjusted regression coefficient_ General vs. non-GDM_ the 3rd trimester_ Random effect model



Sensitivity analysis

S7.5 Figure Adjusted regression coefficients_ exclude studies control for pre-pregnancy BMI or gestational weight gain



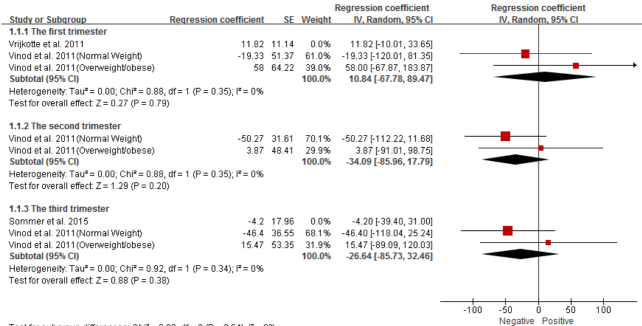
Test for subgroup differences: $Chi^2 = 1.92$, df = 1 (P = 0.17), $I^2 = 47.8\%$

S7.6 Figure Adjusted regression coefficients_exclude studies control for maternal glucose level

				Regression coefficient	Regression coefficient
Study or Subgroup	Regression coefficient	SE	Weight	IV, Random, 95% CI	IV, Random, 95% CI
1.2.1 The first trimes	ter				
Vrijkotte et al. 2011 Subtotal (95% CI)	22.67	9.52	100.0% 100.0%	22.67 [4.01, 41.33] 22.67 [4.01, 41.33]	
Heterogeneity: Not app	olicable				
Test for overall effect:	Z = 2.38 (P = 0.02)				
1.2.2 The second trin	nester				
Crume et al. 2015	17.79	15.1	40.5%	17.79 [-11.81, 47.39]	
Geraghty et al. 2016	27.87	23.35	16.9%	27.87 [-17.90, 73.64]	
Mathews et al. 2003	30.1	14.72	42.6%	30.10 [1.25, 58.95]	
Subtotal (95% CI)			100.0%	24.74 [5.91, 43.57]	•
Heterogeneity: Tau ² =	0.00 ; $Chi^2 = 0.36$, $df = 2$	(P = 0.	83); I ² =	0%	
Test for overall effect:					
1.2.3 The third trimes	ster				
Geraghty et al. 2016	24.85	17.47	28.0%	24.85 [-9.39, 59.09]	-
Mathews et al. 2003	11.1	14.87	38.6%	11.10 [-18.04, 40.24]	
Sommer et al. 2015	-6.1	15.99	33.4%	-6.10 [-37.44, 25.24]	
Ye et al. 2015	9.1	7.91	0.0%	9.10 [-6.40, 24.60]	
Subtotal (95% CI)			100.0%	9.20 [-8.91, 27.31]	-
Heterogeneity: Tau ² =	0.00 ; $Chi^2 = 1.73$, $df = 2$	(P = 0.	42); $I^2 =$	0%	
Test for overall effect:	Z = 1.00 (P = 0.32)	•	-		
					-100 -50 0 5'0 10
Test for subgroup diffe	erences Chi ² = 1.63 df =	2 /P =	∩ 44) l² -	- 0%	Negative Positive

Test for subgroup differences: $Chi^2 = 1.63$, df = 2 (P = 0.44), $I^2 = 0\%$

S7.7 Figure Crude regression coefficients_ exclude studies control for pre-term birth



Test for subgroup differences: $Chi^2 = 0.90$, df = 2 (P = 0.64), $I^2 = 0\%$

S7.8 Figure Adjusted regression coefficients_exclude studies that did not control for pre-term birth

				Regression coefficient	Regression coefficient
Study or Subgroup	Regression coefficient	SE	Weight	IV, Random, 95% CI	IV, Random, 95% CI
1.2.1 The first trimes	ster				
Vrijkotte et al. 2011 Subtotal (95% CI)	22.67	9.52	0.0%	22.67 [4.01, 41.33] Not estimable	
Heterogeneity. Not ap	plicable				
Test for overall effect:	Not applicable				
1.2.2 The second trir	mester				
Crume et al. 2015	17.79	15.1	48.7%	17.79 [-11.81, 47.39]	
Geraghty et al. 2016	27.87	23.35	0.0%	27.87 [-17.90, 73.64]	
Mathews et al. 2003	30.1	14.72	51.3%	30.10 [1.25, 58.95]	
Subtotal (95% CI)			100.0%	24.10 [3.44, 44.76]	•
Heterogeneity: Tau2 =	0.00; Chi ² = 0.34, df = 1	(P = 0.	56); I ² =	0%	
Test for overall effect:	Z = 2.29 (P = 0.02)				
1.2.3 The third trime	ster				
Geraghty et al. 2016	24.85	17.47	0.0%	24.85 [-9.39, 59.09]	
Mathews et al. 2003	11.1	14.87	18.5%	11.10 [-18.04, 40.24]	
Sommer et al. 2015	-6.1	15.99	16.0%	-6.10 [-37.44, 25.24]	- •
Ye et al. 2015	9.1	7.91	65.5%	9.10 [-6.40, 24.60]	- ■
Subtotal (95% CI)			100.0%	7.04 [-5.51, 19.58]	◆
Heterogeneity: Tau2 =	0.00; Chi ² = 0.82, df = 2	(P = 0.	66); I ² =	0%	
Test for overall effect:	Z = 1.10 (P = 0.27)				
					t
					-100 -50 b 5'0 100'
Test for subgroup diff	erences: $Chi^2 = 1.92$ df =	1 (P =	0 17) 12 :	= 47.8%	Negative Positive

Test for subgroup differences: $Chi^2 = 1.92$, df = 1 (P = 0.17), $I^2 = 47.8\%$

High-Density lipoprotein Cholesterol (HDL-C)

S7.3 Table Results summary of the association of maternal HDL-C level with birthweight

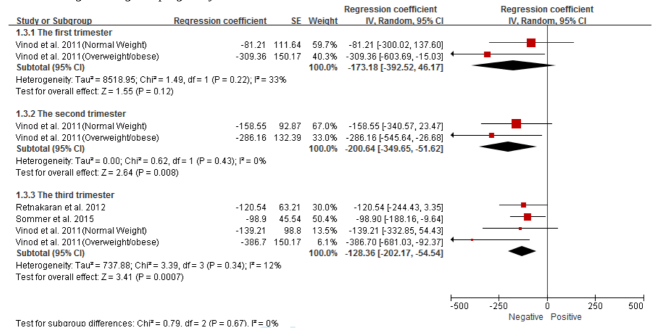
ID	Donulation	Countries	Sample Tri.	Reported	Effect	Lower	Upper	n	Statistical	Orralit	_Th	e co	<u>ntr</u> ol	of co	<u>nf</u> ou	nding	fac	tors
Ш	Population	Countries'	size 171.	measures	size	95%CI	95%CI	p	methods	Quality	a	b	С	d	e	f	g	h
Vinod et al.2011(1)	normal weight	USA	65 1	Crude β	-81.21	-300.02	137.61	ND	SLR	6	×	×	×	×	×	×	×	×
Vinod et al.2011(2)	Overweight/obese	USA	71 1	Crude β	-309.36	-603.69	-15.03	ND	SLR	6	×	×	×	×	×	×	×	×
Wang et al.2015	General	China	636 2	r	-0.12			0.010	Partial correlation	6			×	×	×	×	×	×
Liu et al.2016	General	China	1,546 2	r	-0.01			0.701	Partial correlation	7	×	×	×	×	×	×	×	×
Vinod et al.2011(1)	Normal weight	USA	71 2	Crude β	-158.55	-340.57	23.48	ND	SLR	6	×	×	×	×	×	×	×	×
Vinod et al.2011(2)	Overweight/obese	USA	71 2	Crude β	-286.16	-545.63	-26.68	ND	SLR	6	×	×	×	×	×	×	×	×
Crume et al.2015	General	USA	804 2	Adjusted β	-20.88	-109.69	67.93	0.600) MLR	8			$\sqrt{}$	×	×	×		×
Kulkarni et al.2013	non-GDM	India	631 2	Adjusted β	17.57	-11.64	46.77	ND	MLR	8	×		$\sqrt{}$		×	×		×
Geraghty et al.2016	non-GDM	UK	331 2	Adjusted β	-1236.25	-3322.95	850.45	ND	MLR	7			×			×	×	×
Whyte et al. 2013	General	Ireland	189 2	ND	ND			ND	ND	6	ND	ND	ND	ND	ND	ND	×	ND
Di et al.2005	OGTT+	Italy	83 2	ND	ND			ND	ND	5	ND	ND	ND	ND	ND	ND	×	ND
Zawiejska et al. 2008	GDM	Poland	357 2	ND	ND			ND	ND	5	ND	ND	ND	ND	ND	ND	×	ND
Olmos et al.2014	GDM	Chile	279 2	ND	ND			ND	ND	6	ND	ND	ND	ND	ND	ND	×	ND
Knopp et al.1985	General	USA	248 3	r	-0.06			>0.05	5 Spearman correlati	on 7			×	×	×	×		×
Mossayebi et al.2014	General	Iran	154 3	r	-0.47			<0.00	Pearson correlation	5	×	×	×	×	×	×		×
Charles et al. 2016	General	Multiple	1062 3	r	-0.139			<0.00	Pearson correlation	4	×	×	×	×	×	×	×	×
Vinod et al.2011(1)	Normal weight	USA	69 3	Crude β	-139.21	-332.85	54.43	ND	SLR	6	×	×	×	×	×	×	×	×
Vinod et al.2011(2)	Overweight/obese	USA	70 3	Crude β	-386.70	-681.03	-92.37	ND	SLR	6	×	×	×	×	×	×	×	×
Sommer et al.2015	General	Norway	699 3	Crude β	-98.90	-188.10	-9.60	ND	SLR	9	×	×	×	×	×	×		×
Retnakaran et al.2012	non-GDM	Canada	472 3	Crude β	-120.54	-244.42	3.35	ND	SLR	7	×	×	×	×	×	×		×
Sommer et al.2015	General	Norway	699 3	Adjusted β	-105.40	-183.80	-27.00	ND	MLR	9			$\sqrt{}$	×	×	×		×
Friis et al.2012	General	German	207 3	Adjusted β	-170.00	-329.00	-9.00	0.040	MLR	6		×	×	×	×	×	×	×
Crume et al.2015	General	USA	804 3	Adjusted β	-43.31	-128.33	41.71	0.300) MLR	8			$\sqrt{}$	×	×	×		×
Retnakaran et al.2012	non-GDM	Canada	472 3	Adjusted β	-57.16	-189.42	75.09	ND	MLR	7			$\sqrt{}$					
Kulkarni et al.2013	non-GDM	India	631 3	Adjusted β	-8.89	-38.72	20.95	ND	MLR	8	×		$\sqrt{}$	$\sqrt{}$	×	×		×
Ye et al.2015	non-GDM	China	1,243 3	Adjusted β	-69.50	-110.00	-28.20	ND	MLR	8			$\sqrt{}$					×
Geraghty et al.2016	non-GDM	UK	331 3	Adjusted β	30.00	-114.85	174.84	ND	MLR	7			×			×	×	×
Emet et al.2013	General	Turkey	801 3	p	ND			0.754	Pearson correlation	5	×	×	×	×	×	×	×	×
Couch et al.1998	General	USA	40 3	p	ND			>0.05	5 Pearson correlation	6	×	×	×	×	×	×	×	×
Swierzewska et	General	Poland	136 3	p	ND			>0.05	5 MLR	5	ND	ND	ND	ND	ND	ND	×	ND
Son et al.2010	GDM	Korea	104 3	p	ND			>0.05	5 ND	5	ND	ND	ND	ND	ND	ND		ND
Slagjana et al.2014	non-GDM	Yugoslavia	200 3	ND	ND			ND	ND	5	ND	ND	ND	ND	ND	ND	×	ND
Olmos et al.2014	GDM	Chile	279 3	ND	ND			ND	ND	6	ND	ND	ND	ND	ND	ND	×	ND

The bold font represents statistically significant results. r: Correlation coefficients; β : regression coefficients.

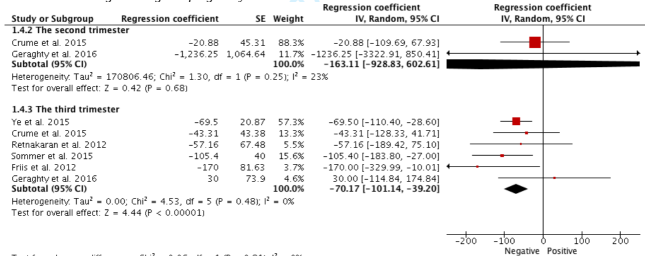
Confounding factors: a. Gestational age; b. Neonatal gender; c. Maternal age; d. Pre-pregnancy BMI; e. Gestational weight gain; f. Maternal glucose level; g. pre-term birth; h. Maternal lipid levels. Abbreviation: Trimesters(Tri.), Gestational diabetes mellitus(GDM), Positive screenes of Oral Glucose Tolerance test(OGTT+), Confidence interval(CI), Not documented(ND), Simple linear regression(SLR), Multiple linear regression(MLR), United Kingdom(UK).

Meta-analysis

S7.9 Figure Overall meta-analysis of crude regression coefficients for the association between maternal HDL-C levels and birthweight throughout pregnancy



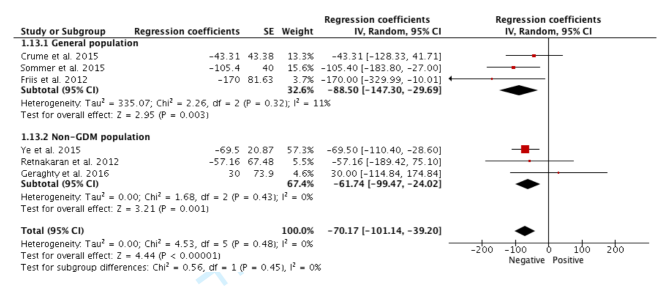
S7.10 Figure Overall meta-analysis of adjusted regression coefficients for the association between maternal HDL-C levels and birthweight throughout pregnancy



Test for subgroup differences: $Chi^2 = 0.06$, df = 1 (P = 0.81), $I^2 = 0\%$

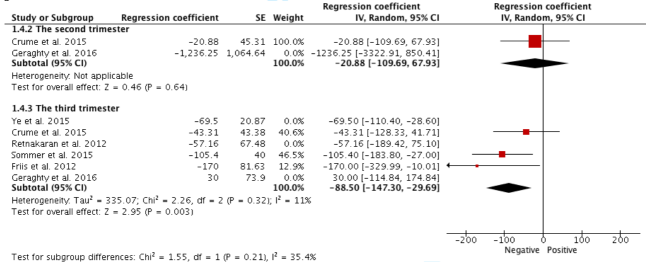
Subgroup analysis

S7.11 Figure Adjusted regression coefficient_ General vs. non-GDM_ the 3rd trimester_ Random effect model

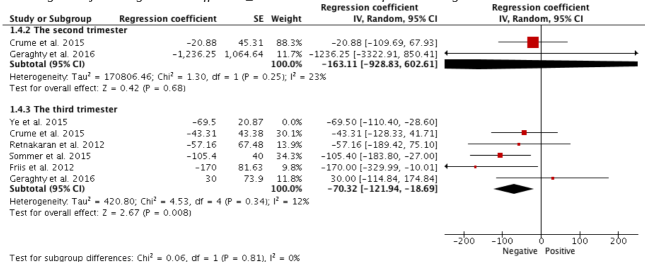


Sensitivity analysis

S7.12 Figure Adjusted regression coefficients_ exclude studies control for pre-pregnancy BMI or gestational weight gain



S7.13 Figure Adjusted regression coefficients exclude studies control for maternal glucose level



S7.14 Figure Adjusted regression coefficients_ exclude studies control for pre-term birth Regression coefficient

				Regression coefficient	Kegr	ession coefficient	
Study or Subgroup	Regression coefficient	SE	Weight	IV, Random, 95% CI	IV,	Random, 95% CI	
1.4.2 The second trime	ester						
Crume et al. 2015	-20.88	45.31	88.3%	-20.88 [-109.69, 67.93]	_		
Geraghty et al. 2016 Subtotal (95% CI)	-1,236.25	1,064.64	11.7% 100.0%	-1236.25 [-3322.91, 850.41] -163.11 [-928.83, 602.61]			
Heterogeneity: Tau ² = 1	170806.46; Chi² = 1.30, df	r = 1 (P = 0)).25); I ² =	= 23%			
Test for overall effect: Z	= 0.42 (P = 0.68)						
a a 2 The ability selection							
1.4.3 The third trimest	ter						
Ye et al. 2015	-69.5	20.87	53.0%	-69.50 [-110.40, -28.60]		-	
Crume et al. 2015	-43.31	43.38	16.6%	-43.31 [-128.33, 41.71]			
Retnakaran et al. 2012	-57.16	67.48	0.0%	-57.16 [-189.42, 75.10]			
Sommer et al. 2015	-105.4	40	19.2%	-105.40 [-183.80, -27.00]			
Friis et al. 2012	-170	81.63	5.1%	-170.00 [-329.99, -10.01]			
Geraghty et al. 2016	30	73.9	6.1%	30.00 [-114.84, 174.84]		•	-
Subtotal (95% CI)			100.0%	-71.02 [-107.65, -34.38]	•		
Heterogeneity: Tau ² = 2	223.11; Chi ² = 4.49, df = 4	1 (P = 0.34)	$1: I^2 = 11:$	%			
Test for overall effect: Z			,				
							-1-
					-2'00 -1'00		200
	51.2 0.05 16 4		17 001		N e	egative Positive	

Test for subgroup differences: $\mathrm{Chi^2} = 0.06$, $\mathrm{df} = 1$ (P = 0.81), $\mathrm{I^2} = 0\%$

Low-Density lipoprotein Cholesterol (LDL-C)

S7.4 Table Results summary of the association of maternal LDL-C level with birthweight

ID	Population	Countries	Sample Tri	Reported		Lower		n	Statistical	Quality	Th	ie co	ntro	ol of c	onfor	ınding	fact	ors
10	1 opulation	Countries	size 111	· measures	size	95%CI	95%CI	р	methods	Quanty	A	b	c	d	e	f	g	h
Vinod et al.2011(1)	Normal weight	USA	65 1	Crude β	-34.80	-152.92	83.32	ND	SLR	6	×	×	×	×	×	×	×	×
Vinod et al.2011(2)	Overweight/obese	USA	71 1	Crude β	108.28	-42.76	259.31	ND	SLR	6	×	×	×	×	×	×	×	×
Liu et al.2016	General	China	1,546 2	r	-0.01			0.843	Partial correlation	7	×	×	×	×	×	×	×	×
Vinod et al.2011(1)	Normal weight	USA	71 2	Crude β	-58.00	-133.52	17.51	ND	SLR	6	×	×	×	×	×	×	×	×
Vinod et al.2011(2)	Overweight/obese	USA	71 2	Crude β	34.80	-83.32	152.92	ND	SLR	6	×	×	×	×	×	×	×	×
Geraghty et al.2016	non-GDM	UK	331 2	Adjusted β	18.39	-38.44	75.21	ND	MLR	7			×	\checkmark		×	×	×
Wang et al.2015	General	China	636 2	ND	ND			ND	Partial correlation	6			×	×	×	×	×	×
Whyte et al. 2013	General	Ireland	189 2	ND	ND			ND	ND	6	ND	ND	N	ND	ND	ND	×	ND
Di et al.2005	OGTT+	Italy	83 2	ND	ND			ND	ND	5	ND	ND	N	ND	ND	ND		ND
Olmos et al.2014	GDM	Chile	279 2	ND	ND			ND	ND	6	ND	ND	N	ND	ND	ND	×	ND
Knopp et al.1985	General	USA	248 3	r	0.01			>0.05	Spearman correlation	n 7			×	×	×	×		×
Mossayebi et al.2014	General	Iran	154 3	r	0.40			< 0.001	Pearson correlation	5	×	×	×	×	×	×		×
Charles et al. 2016	General	Multiple	1062 3	r	0.001			<0.000	Pearson correlation	n 4	×	×	×	×	×	×	×	×
Retnakaran et al.2012	non-GDM	Canada	472 3	Crude β	-15.22	-55.49	25.05	ND	SLR	7	×	×	×	×	×	×		×
Vinod et al.2011(1)	Normal weight	USA	69 3	Crude β	-50.27	-131.60	31.06	ND	SLR	6	×	×	×	×	×	×	×	×
Vinod et al.2011(2)	Overweight/obese	USA	70 3	Crude β	38.67	-79.45	156.79	ND	SLR	6	×	×	×	×	×	×	×	×
Ye et al.2015	non-GDM	China	1,243 3	Adjusted β	35.40	10.10	60.80	ND	MLR	8								×
Retnakaran et al.2012	non-GDM	Canada	472 3	Adjusted β	-6.79	-46.98	33.39	ND	MLR	7						$\sqrt{}$		$\sqrt{}$
Geraghty et al.2016	non-GDM	UK	331 3	Adjusted β	19.97	-24.34	64.27	ND	MLR	7			×			×	×	×
Emet et al.2013	General	Turkey	801 3	p	ND			0.440	Pearson correlation	5	×	×	×	×	×	×	×	×
Couch et al.1998	General	USA	40 3	p	ND			>0.05	Pearson correlation	6	×	×	×	×	×	×	×	×
Swierzewska et al.2015	General	Poland	136 3	p	ND			>0.05	MLR	5	ND	ND	N	ND	ND	ND	×	ND
Sommer et al.2015	General	Norway	699 3	ND	ND			ND	ND	9	ND	ND	N	ND	ND	ND	×	ND
Slagjana et al.2014	non-GDM	Yugoslavia	200 3	ND	ND			ND	ND	5	ND	ND	N	ND	ND	ND	X	ND
Son et al.2010	GDM	Korea	104 3	ND	ND			ND	ND	5	ND	ND	N	ND	ND	ND		ND
Olmos et al.2014	GDM	Chile	279 3	ND	ND			ND	ND	6	ND	ND	N	ND	ND	ND	X	ND

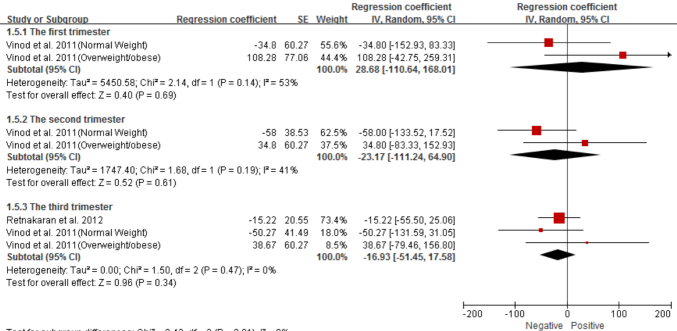
The bold font represents statistically significant results.

r: Correlation coefficients; β : regression coefficients.

Confounding factors: a. Gestational age; b. Neonatal gender; c. Maternal age; d. Pre-pregnancy BMI; e. Gestational weight gain; f. Maternal glucose level; g. pre-term birth; h. Maternal lipid levels. Abbreviation: Trimesters(Tri.), Gestational diabetes mellitus(GDM), Positive screenes of Oral Glucose Tolerance test(OGTT+), Confidence interval(CI), Not documented(ND), Simple linear regression(SLR), Multiple linear regression(MLR), United Kingdom(UK).

Meta-analysis

S7.15 Figure Overall meta-analysis of crude regression coefficients for the association between maternal LDL-C levels and birthweight throughout pregnancy



Test for subgroup differences: $Chi^2 = 0.42$, df = 2 (P = 0.81), $I^2 = 0\%$

S7.16 Figure Overall meta-analysis of adjusted regression coefficients for the association between maternal LDL-C levels and birthweight throughout pregnancy

0 0	1 0						
			1	Regression coefficient		Regression coefficient	
Study or Subgroup	Regression coefficient	SE	Weight	IV, Random, 95% CI		IV, Random, 95% CI	
1.6.2 The second trimes	ster					<u>_</u>	
Geraghty et al. 2016	18.39	28.99	100.0%	18.39 [-38.43, 75.21]			_
Subtotal (95% CI)			100.0%	18.39 [-38.43, 75.21]			_
Heterogeneity: Not appli	icable						
Test for overall effect: Z	= 0.63 (P = 0.53)						
1.6.3 The third trimeste	er						
Ye et al. 2015	35.4	12.93	48.2%	35.40 [10.06, 60.74]			
Retnakaran et al. 2012	-6.79	20.5	27.7%	-6.79 [-46.97, 33.39]			
Geraghty et al. 2016	19.97	22.6	24.1%	19.97 [-24.33, 64.27]			
Subtotal (95% CI)			100.0%	19.98 [-5.25, 45.20]			
Heterogeneity: Tau ² = 17	76.80; Chi²= 3.05, df= 2 (P	= 0.22)	; I² = 34%				
Test for overall effect: Z =	= 1.55 (P = 0.12)						
					-100	-50 0 50	100
Toot for outbaroup differe	oncoc: Chi3 = 0.00 df = 1./E	0.00	N IZ — O OV			Negative Positive	

Test for subaroup differences: Chi² = 0.00, df = 1 (P = 0.96), l² = 0%

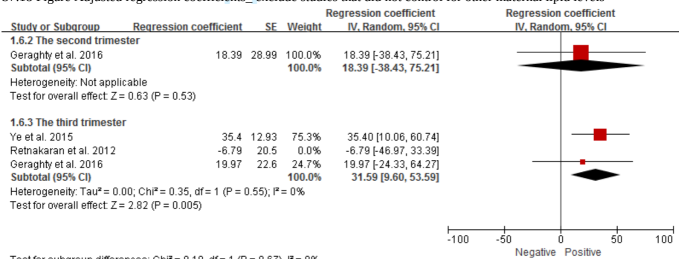
Sensitivity analysis

S7.17 Figure Adjusted regression coefficients_ exclude studies that did not control for pre-term birth

Study or Subgroup	Dograpaion coefficient	er.		Regression coefficient		_	on coefficient	
Study or Subgroup	Regression coefficient	3E	Weight	IV, Random, 95% CI		IV, Kalic	lom, 95% CI	
1.6.2 The second trime								
Geraghty et al. 2016	18.39	28.99	0.0%	18.39 [-38.43, 75.21]				
Subtotal (95% CI)				Not estimable				
Heterogeneity: Not app	licable							
Test for overall effect: N	lot applicable							
1.6.3 The third trimest	er							
Ye et al. 2015	35.4	12.93	57.1%	35.40 [10.06, 60.74]				
Retnakaran et al. 2012	-6.79	20.5	42.9%	-6.79 [-46.97, 33.39]				
Geraghty et al. 2016	19.97	22.6	0.0%	19.97 [-24.33, 64.27]				
Subtotal (95% CI)			100.0%	17.30 [-23.62, 58.23]				
Heterogeneity: Tau ² = 5	596.28; Chi ² = 3.03, df = 1 (P	= 0.08)	: I² = 67%					
Test for overall effect: Z	(= 0.83 (P = 0.41)	,						
					100	- 	 	400
					-100	-50	0 50	100
T16						Negativ	e Positive	

Test for subgroup differences: Not applicable

S7.18 Figure Adjusted regression coefficients_ exclude studies that did not control for other maternal lipid levels



Test for subgroup differences: Chi² = 0.18, df = 1 (P = 0.67), l² = 0%

Triglycerides (TG)

S7.5 Table Results summary of the association of maternal TG level with birthweight

ID	ID Population Countries Sample Size Tri. Reported Effect Lower Upper p Statistical methods are size 95%CI 95%CI p Statistical methods.	Statistical methods	Onality		he (cont		of co		und	ling							
12	1 opulation	Countries	size	measures	size	95%CI	95%CI	Р	Statistical methods	Quanty		b	c				g	h
Nolan et al.1995	General	Australia	388 1	r	0.12			0.020	Univariate analyses	6			×	×	×	×	×	×
Vinod et al.2011(1)	Normal weight	USA	65 1	Crude β	132.86	13.11	252.62	ND	SLR	6	×	×	×	×	×	×	×	×
Vinod et al.2011(2)	Overweight/obese	USA	71 1	Crude β	124.00	-40.10	288.11	ND	SLR	6	×	×	×	×	×	×	×	×
Vrijkotte et al.2011	General	Netherlands	2,052 1	Crude β	47.14	12.42	81.87	ND	Univariate analyses	8			×	×	×	×	$\sqrt{}$	×
Vrijkotte et al.2011	General	Netherlands	2,052 1	Adjusted β	86.72	56.13	117.30	ND	MLR	8			$\sqrt{}$			×	$\sqrt{}$	×
Harmon et al.2011	non-GDM	USA	38 1	p	ND			>0.05	Pearson correlation	5	×	×	×	×	×	×	×	×
Liu et al.2016	General	China	1,546 2	r	0.10			<0.001	Partial correlation	7	×	×	×	×	×	×	×	×
Wang et al.2015	General	China	636 2	r	0.19			< 0.01	Partial correlation	6			×	×	×	×	×	×
Di et al.2005	OGTT+	Italy	83 2	r	0.30			< 0.05	SLR	5	×	×	×	×	×	×	×	×
Zawiejska et al. 2008	GDM	Poland	357 2	r	0.14			<0.01	SLR	5	×	×	×	×	×	×	×	×
Vinod et al.2011(1)	Normal weight	USA	71 2	Crude β	97.43	4.29	190.57	ND	SLR	6	×	×	×	×	×	×	×	×
Vinod et al.2011(2)	Overweight/obese	USA	71 2	Crude β	132.86	4.24	261.49	ND	SLR	6	×	×	×	×	×	×	×	×
Crume et al.2015	General	USA	804 2	Adjusted β	7.97	-44.19	60.13	0.700	MLR	8				×	×	×		×
Kulkarni et al.2013	non-GDM	India	631 2	Adjusted β	14.76	-13.34	42.86	ND	MLR	8	×		$\sqrt{}$		×	×	$\sqrt{}$	×
Hwang et al.2015	non-GDM	Korea	1,011 2	Adjusted β^	7125.42	1693.49	12557.35	0.002	MLR	8			$\sqrt{}$	×		×	×	×
Whyte et al. 2013	General	Ireland	189 2	p	+			< 0.05	SLR	6	×	×	×	×	×	×	×	×
Geraghty et al.2016	non-GDM	UK	331 2	p	ND			>0.1	MLR	7			×			×	×	×
Olmos et al.2014	GDM	Chile	279 2	ND	ND			ND	ND	6	ND	ND	ND	ND	ND	ND	×	ND
Mossayebi et al.2014	General	Iran	154 3	r	0.68			<0.001	Pearson correlation	5	×	×	×	×	×	×	$\sqrt{}$	×
Charles et al. 2016	General	Multiple	1062 3	r	-0.014			<0.000	1Pearson correlation	4	×	×	×	×	×	×	×	×
Son et al.2010	GDM	Korea	104 3	r	0.17			0.070	ND	5	×	×	×	×	×	×	$\sqrt{}$	×
Ahmad et al. 2006	non-GDM	Malaysia	246 3	r	0.12			0.057	Univariate analyses	6		×	×	×	×	×	$\sqrt{}$	×
Couch et al.1998(1)	non-GDM	USA	20 3	r	0.46			< 0.05	Pearson correlation	6	×	×	×	×	×	×	×	×
Slagjana et al.2014	non-GDM	Yugoslavia	200 3	r	0.16			0.077	Correlation analysis	5	×	×	×	×	×	×	×	×
Olmos et al.2014(1)	GDM-normal weight	Chile	128 3	r	0.12			0.158	SLR	6	×	×	×	×	×	×	×	×
Olmos et al.2014(2)	GDM-overweight	Chile	105 3	r	0.42			<0.001	I SLR	6	×	×	×	×	×	×	×	×
Olmos et al.2014(3)	GDM-obese	Chile	46 3	r	0.47			<0.001	I SLR	6	×	×	×	×	×	×	×	×
Kitajima et al.2001	OGTT +	Japan	146 3	r	0.22			0.009	SLR	6	×	×	×	×	×	×	$\sqrt{}$	×

ID	Population	Countries	Sample Tri.	Reported	Effect	Lower	Upper	р	Statistical methods	Ouality		e co		l of act		four	nding
10	Topulation	Countries	size	measures	size	95%CI	95%CI	Р	Statistical methods	Quanty		b c				f g	h
Knopp et al.1992(1)	OGTT-	USA	521 3	r	0.09			≤0.05	Spearman correlation	6	×	× ×	×		× :	× ×	×
Knopp et al.1992(2)	OGTT+ plus GDM	USA	264 3	r	0.16			≤0.01	Spearman correlation	6	×	× ×	×		×	× ×	×
Vinod et al.2011(1)	Normal weight	USA	69 3	Crude β	79.72	-8.99	168.42	ND	SLR	6	×	× ×	×		× :	× ×	×
Vinod et al.2011(2)	Overweight/obese	USA	70 3	Crude β	168.29	52.97	283.61	ND	SLR	6	×	× ×	×		× :	× ×	×
Sommer et al.2015	General	Norway	699 3	Crude β	48.80	-14.80	112.40	ND	SLR	9	×	× ×	×		×	× √	×
Retnakaran et al.2012	non-GDM	Canada	472 3	Crude β	61.11	-1.18	123.40	ND	SLR	7	×	× ×	×		× :	× √	×
Sommer et al.2015	General	Norway	699 3	Adjusted β	94.40	37.80	150.90	ND	MLR	9	$\sqrt{}$	√ \	l ×		×	× √	×
Retnakaran et al.2012	non-GDM	Canada	472 3	Adjusted β	-1.59	-70.67	67.49	ND	MLR	7	$\sqrt{}$	√ \	/ √	1	√ .	V V	
Brunner et al.2013	General	German	208 3	Adjusted β	-47.83	-138.75	43.09	>0.05	MLR	7	$\sqrt{}$	√ >	< √		√ .	V ×	×
Friis et al.2012	General	German	207 3	Adjusted β	94.00	2.00	187.00	0.046	MLR	6	$\sqrt{}$	× ×	×		× :	× ×	×
Mossayebi et al.2014	General	Iran	154 3	Adjusted β	464.13	370.24	558.02	ND	MLR	5	×	√ >	×		×	× √	×
Crume et al.2015	General	USA	804 3	Adjusted β	17.71	-24.01	59.44	0.400	MLR	8	$\sqrt{}$	√ \	/ ×		× :	× √	×
Geraghty et al.2016	non-GDM	UK	331 3	Adjusted β	111.18	8.48	213.87	ND	MLR	7	$\sqrt{}$	√ >	< √		√ :	× ×	×
Ye et al.2015	non-GDM	China	1,243 3	Adjusted β	25.20	7.90	42.60	ND	MLR	8	$\sqrt{}$	√ \	1 1	1	√ .	V V	×
Kulkarni et al.2013	non-GDM	India	631 3	Adjusted β	36.27	4.32	68.23	ND	MLR	8	×	√ ₁	1 1		× :	× √	×
Hwang et al.2015	non-GDM	Korea	1,011 3	Adjusted β^{\wedge}	11609.12	6177.20	17041.05	<0.000	1MLR	8	$\sqrt{}$	√ \	/ ×		√ :	× ×	×
Swierzewska et al.2015	General	Poland	136 3	p	ND			>0.05	MLR	5	NDN	IDN	D NI	D N	NDN	D ×	ND
Emet et al.2013	General	Turkey	801 3	$p\P$	+			0.033	Pearson correlation	5	×	× ×	×		×	××	×
Schaefer-Graf et al.2011	non-GDM	German	190 3	p	ND			>0.05	Pearson correlation	5	×	× ×	×		×	× √	×
Couch et al.1998(2)	GDM	USA	20 3	p	ND			>0.05	Pearson correlation	6	×	× ×	×		×	××	×
Schaefer-Graf et al.2008	GDM	German	150 3	р	ND			>0.05	Spearman correlation	5	×	× ×	×		×	××	×

The bold font represents statistically significant results.

Confounding factors: a. Gestational age; b. Neonatal gender; c. Maternal age; d. Pre-pregnancy BMI; e. Gestational weight gain; f. Maternal glucose level; g. pre-term birth; h. Maternal lipid levels. Abbreviation: Trimesters(Tri.), Gestational diabetes mellitus(GDM), Positive screenes of Oral Glucose Tolerance test(OGTT+), Confidence interval(CI), Not documented(ND), Simple linear regression(SLR), Multiple linear regression(MLR), United Kingdom(UK).

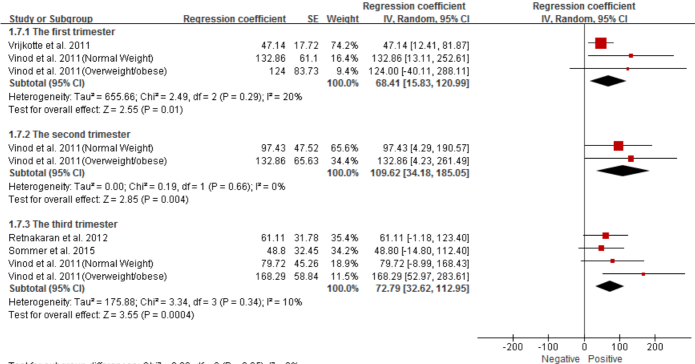
[^] Maternal TG level was log-transformed

[¶] Exposure of this study is change in maternal TG level from the first trimester to the third trimester

r: Correlation coefficients; β: regression coefficients.

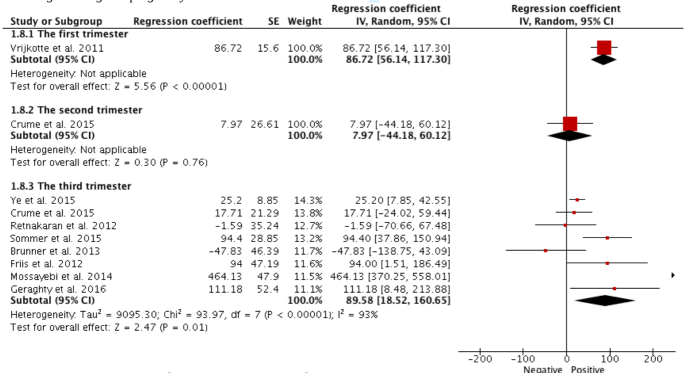
Meta-analysis

S7.19 Figure Overall meta-analysis of crude regression coefficients for the association between maternal TG levels and birthweight throughout pregnancy



Test for subgroup differences: $Chi^2 = 0.86$, df = 2 (P = 0.65), $I^2 = 0\%$

S7.20 Figure Overall meta-analysis of adjusted regression coefficients for the association between maternal TG levels and birthweight throughout pregnancy



Test for subgroup differences: $Chi^2 = 6.87$, df = 2 (P = 0.03), $I^2 = 70.9\%$

Subgroup analysis

S7.21 Figure Adjusted regression coefficient_ General vs. non-GDM_ the 3rd trimester_ Random effect model

				Regression coefficient	Regression coefficient
Study or Subgroup	Regression coefficient	SE	Weight	IV, Random, 95% CI	IV, Random, 95% CI
1.14.1 General populat	tion				
Crume et al. 2015	17.71	21.29	13.8%	17.71 [-24.02, 59.44]	 -
Sommer et al. 2015	94.4	28.85	13.2%	94.40 [37.86, 150.94]	
Brunner et al. 2013	-47.83	46.39	11.7%	-47.83 [-138.75, 43.09]	
Friis et al. 2012	94	47.19	11.6%	94.00 [1.51, 186.49]	-
Mossayebi et al. 2014	464.13	47.9		464.13 [370.25, 558.01]	
Subtotal (95% CI)			61.9%	122.31 [-15.77, 260.39]	
Heterogeneity: Tau ² = 2	?3250.17; Chi² = 80.79, df	= 4 (P)	< 0.0000)1); I ² = 95%	
Test for overall effect: Z	= 1.74 (P = 0.08)				
1.14.2 Non-GDM popu	ılation				
Ye et al. 2015		8.85	14.3%	25.20 [7.85, 42.55]	+
Retnakaran et al. 2012		35.24			
Geraghty et al. 2016	111.18	52.4	11.1%		
Subtotal (95% CI)			38.1%		◆
Heterogeneity: Tau ² = 5	78.32; Chi ² = 3.26, df = 2	P = 0.	$20); I^2 =$	39%	
Test for overall effect: Z		,			
Total (95% CI)			100.0%	89.58 [18.52, 160.65]	•
Heterogeneity: Tau ² = 9	9095.30; Chi ² = 93.97, df :	= 7 (P <	0.00001	L); I ² = 93%	-200 0 100 200
Test for overall effect: Z					Negative Positive
Test for subgroup differ	rences: Chi² = 1.59, df <u>=</u> 1	$(P_{-} = 0.2)$	$(2.1), 1^2 = 3$	37.0%	regative rositive

Sensitivity analysis

S7.22 Figure Adjusted regression coefficients_ the 3rd trimester_ exclude studies control for pre-pregnancy BMI or gestational weight gain

				Regression coefficient	Regression coefficient
Study or Subgroup	Regression coefficient	SE	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Crume et al. 2015	17.71	21.29	25.9%	17.71 [-24.02, 59.44]	
Sommer et al. 2015	94.4	28.85	25.6%	94.40 [37.86, 150.94]	_
Brunner et al. 2013	-47.83	46.39	0.0%	-47.83 [-138.75, 43.09]	
Friis et al. 2012	94	47.19	24.3%	94.00 [1.51, 186.49]	-
Mossayebi et al. 2014	464.13	47.9	24.2%	464.13 [370.25, 558.01]	·
Ye et al. 2015	25.2	8.85	0.0%	25.20 [7.85, 42.55]	
Retnakaran et al. 2012	-1.59	35.24	0.0%	-1.59 [-70.66, 67.48]	
Geraghty et al. 2016	111.18	52.4	0.0%	111.18 [8.48, 213.88]	
Total (95% CI)			100.0%	163.95 [3.26, 324.65]	
Heterogeneity: Tau ² = 25 Test for overall effect: Z =	5461.60; Chi ² = 72.54, dt = 2.00 (P = 0.05)	f = 3 (P	< 0.0000	01); I² = 96%	-200 -100 0 100 200 Negative Positive

S7.23 Figure Adjusted regression coefficients_ the 3rd trimester_ exclude studies control for maternal glucose level

				Regression coefficient	Regression coefficient
Study or Subgroup	Regression coefficient	SE	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Crume et al. 2015	17.71	21.29	21.1%	17.71 [-24.02, 59.44]	
Sommer et al. 2015	94.4	28.85	20.8%	94.40 [37.86, 150.94]	_
Brunner et al. 2013	-47.83	46.39	0.0%	-47.83 [-138.75, 43.09]	
Friis et al. 2012	94	47.19	19.5%	94.00 [1.51, 186.49]	-
Mossayebi et al. 2014	464.13	47.9	19.5%	464.13 [370.25, 558.01]	·
Ye et al. 2015	25.2	8.85	0.0%	25.20 [7.85, 42.55]	
Retnakaran et al. 2012	-1.59	35.24	0.0%	-1.59 [-70.66, 67.48]	
Geraghty et al. 2016	111.18	52.4	19.1%	111.18 [8.48, 213.88]	
Total (95% CI)			100.0%	153.36 [19.84, 286.89]	
Heterogeneity. Tau ² = 21 Test for overall effect: Z =	L529.40; Chi ^z = 72.66, di = 2.25 (P = 0.02)	f = 4 (P	< 0.0000)1); I ² = 94%	-200 -100 0 100 200 Negative Positive

S7.24 Figure Adjusted regression coefficients_ the 3rd trimester_ exclude studies control for other maternal lipid levels

Regression coefficient

Regression coefficient

				Regression coefficient	Regression coefficient
Study or Subgroup	Regression coefficient	SE	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Crume et al. 2015	17.71	21.29	15.6%	17.71 [-24.02, 59.44]	
Sommer et al. 2015	94.4	28.85	15.1%	94.40 [37.86, 150.94]	
Brunner et al. 2013	-47.83	46.39	13.5%	-47.83 [-138.75, 43.09]	
Friis et al. 2012	94	47.19	13.4%	94.00 [1.51, 186.49]	•
Mossayebi et al. 2014	464.13	47.9	13.3%	464.13 [370.25, 558.01]	·
Ye et al. 2015	25.2	8.85	16.2%	25.20 [7.85, 42.55]	-
Retnakaran et al. 2012	-1.59	35.24	0.0%	-1.59 [-70.66, 67.48]	
Geraghty et al. 2016	111.18	52.4	12.9%	111.18 [8.48, 213.88]	
Total (95% CI)			100.0%	103.46 [23.05, 183.88]	-
Heterogeneity: Tau ² = 1	0326.69; Chi ^z = 92.56, d	f = 6 (P	< 0.0000	01); I ² = 94%	-200 -100 0 100 200
Test for overall effect: Z	= 2.52 (P = 0.01)				Negative Positive

S7.25 Figure Adjusted regression coefficients_ the 3rd trimester_ exclude studies control for pre-term birth

				Regression coefficient	Regression coefficient
Study or Subgroup	Regression coefficient	SE	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Crume et al. 2015	17.71	21.29	0.0%	17.71 [-24.02, 59.44]	
Sommer et al. 2015	94.4	28.85	0.0%	94.40 [37.86, 150.94]	
Brunner et al. 2013	-47.83	46.39	34.3%	-47.83 [-138.75, 43.09]	
Friis et al. 2012	94	47.19	33.9%	94.00 [1.51, 186.49]	-
Mossayebi et al. 2014	464.13	47.9	0.0%	464.13 [370.25, 558.01]	
Ye et al. 2015	25.2	8.85	0.0%	25.20 [7.85, 42.55]	
Retnakaran et al. 2012	-1.59	35.24	0.0%	-1.59 [-70.66, 67.48]	
Geraghty et al. 2016	111.18	52.4	31.8%	111.18 [8.48, 213.88]	
Total (95% CI)			100.0%	50.87 [-49.57, 151.30]	
Heterogeneity: Tau² = 55 Test for overall effect: Z =	, ,	2 (P = 1	0.04); l ² =	= 70%	-200 -100 0 100 200 Negative Positive

S7.26 Figure Adjusted regression coefficients_ the 3rd trimester_ exclude studies that did not control for gestational age

				Regression coefficient	Regression coefficient
Study or Subgroup	Regression coefficient	SE	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Crume et al. 2015	17.71	21.29	19.9%	17.71 [-24.02, 59.44]	
Sommer et al. 2015	94.4	28.85	15.4%	94.40 [37.86, 150.94]	_
Brunner et al. 2013	-47.83	46.39	8.6%	-47.83 [-138.75, 43.09]	
Friis et al. 2012	94	47.19	8.4%	94.00 [1.51, 186.49]	-
Mossayebi et al. 2014	464.13	47.9	0.0%	464.13 [370.25, 558.01]	
Ye et al. 2015	25.2	8.85	28.1%	25.20 [7.85, 42.55]	-
Retnakaran et al. 2012	-1.59	35.24	12.4%	-1.59 [-70.66, 67.48]	- + -
Geraghty et al. 2016	111.18	52.4	7.2%	111.18 [8.48, 213.88]	_
Total (95% CI)			100.0%	36.72 [5.29, 68.14]	•
Heterogeneity: $Tau^2 = 83$	36.05; Chi ² = 13.46, df =	6 (P = 1	0.04); l2 :	= 55%	
Test for overall effect: Z	= 2.29 (P = 0.02)				-2'00 -1'00 Ó 100 200

Free Fatty Acids (FFAs)

S7.6 Table Results summary of the association of maternal FFAs levels with birthweight

ID	Donulation	Population Countries		Countries	opulation Countries Sample Tri. Reported Effect Lower Upper p Statistical methods Quali		Ouglity	The	con	trol o	f cor	ıfour	nding	g fact	ors	FFAs'			
ID	1 opulation	Countries	size	111.	measures	size	95%CI 9	5%CI	p Statistical methods	Quanty	a	b	c	d	e	f	g	h	unit
Harmon et al.2011	non-GDM	USA	38	1	p	ND			>0.05 Pearson correlation	5	×	×	×	×	×	×	×	×	$\mu E q/L$
Crume et al.2015	General	USA	804	2	Adjusted β	0.06	-0.12	0.24	0.500 MLR	8			$\sqrt{}$	×	×	×	$\sqrt{}$	×	mg/dL
Crume et al.2015	General	USA	804	3	Adjusted β	0.21	0.01	0.41	0.030 MLR	8				×	×	×		×	mg/dL
Knopp et al.1985	General	USA	248	3	r	0.002			>0.05 Spearman correlation	7			×	×	×	×	$\sqrt{}$	×	μmol/L
Kitajima et al.2001	OGTT +	Japan	146	3	r	0.03			0.730 SLR	6	×	×	×	×	×	×		× 1	mEq/dL
Schaefer-Graf et al.2008	GDM	German	150	3	r	0.27			0.002 Spearman correlation	5	×	×	×	×	×	×	×	×	μmol/L
Couch et al.1998	General	USA	40	3	p	ND			>0.05 Pearson correlation	6	×	×	×	×	×	×	×	×	mg/dL
Friis et al.2012	General	German	207	3	p	ND			>0.05 MLR	6	$\sqrt{}$	×	×	×	×	×	×	×	ND
Schaefer-Graf et al.2011	non-GDM	German	190	3	p	ND			>0.05 Pearson correlation	5	×	×	×	×	×	×		×	μmol/L

The bold font represents statistically significant results.

Confounding factors: a. Gestational age; b. Neonatal gender; c. Maternal age; d. Pre-pregnancy BMI; e. Gestational weight gain; f. Maternal glucose level; g. pre-term birth; h. Maternal lipid levels.

Abbreviation: Trimesters(Tri.), Gestational diabetes mellitus(GDM), Positive screenes of Oral Glucose Tolerance test(OGTT+), Confidence interval(CI), Not documented(ND), Simple linear regression(SLR), Multiple linear regression(MLR).

Very Low-density lipoprotein cholesterol (VLDL)

S7.7 Table Results summary of the association of maternal VLDL-C levels with birthweight

ID	Population	Countries	Sample Trimester		Reported	Effect	C4 - 42 - 42 1 41 1	01'4	The control of confounding factors								
			size	1 rimester	measures	size p	p Statistical methods	Quanty	a	b	c	d	e	f	g	h	
Couch et al.1998	General	USA	40	3	p	ND	>0.05 Pearson correlation	6	×	×	×	×	×	×	×	×	
Knopp et al.1985	General	USA	248	3	r	0.03	>0.05 Spearman correlation	7	$\sqrt{}$	$\sqrt{}$	×	×	×	×		×	

r: Correlation coefficients

Confounding factors: a. Gestational age; b. Neonatal gender; c. Maternal age; d. Pre-pregnancy BMI; e. Gestational weight gain; f. Maternal glucose level; g. pre-term birth; h. Maternal lipid levels. Abbreviation: Not documented(ND).

r: Correlation coefficients; β: regression coefficients.

Supplementary 8 Data analysis for Large for gestational age

Total cholesterol (TC)

S8.1 Table Results summary of the association of maternal TC levels with LGA

Study ID	Population		Sample	Trimesters	Reported		Lower		р	Statistical methods	Quality		e con		f conf tors	ding	
	-		size		measures	size	95%CI	95%CI				a	b	c	d	e	f
Jin et al.2016	non-GDM	China	934	1	ND	ND)		ND	ND	7	ND	ND	ND	ND	×	ND
Vrijkotte et al.2012	non-GDM	Netherlands	4,008	1	Crude OR	1.10	0.97	1.25	ND	Logistic regression	8	×	×	×	×	×	×
Vrijkotte et al.2012	non-GDM	Netherlands	4,008	1	Adjusted O	R 1.08	0.95	1.22	ND	MLOR	8		\checkmark	×	×	×	×
Jin et al.2016	non-GDM	China	934	2	ND	ND)		ND	ND	7	ND	ND	ND	ND		ND
Di et al.2005	OGTT+	Italy	83	2	ND	ND)		ND	ND	5	ND	ND	ND	ND	×	ND
Mossayebi et al.2014	General	Iran	82	3	Crude OR	* 13.30	2.80	62.50	ND	Chi-squared test	5	×	×	X	×		×
Mossayebi et al.2014	General	Iran	82	3	Adjusted OI	R* 1.10	0.20	8.10	ND	MLOR	5			×	$\sqrt{}$		
Ye et al.2015	non-GDM	China	1,204	3	Adjusted O	R 1.04	0.94	1.15	ND	MLOR	8				$\sqrt{}$		×
Jin et al.2016	non-GDM	China	934	3	Adjusted O	R 0.98	0.81	1.11	0.715	MLOR	7				×		×
Hou et al.2014	non-GDM	China	2,790	3	Adjusted OI	R¶ 1.08	0.75	1.56	ND	MLOR	7			×	×		×
Schaefer-Graf et al.2008	GDM	German	150	3	p	ND			>0.05	MLOR	5				$\sqrt{}$	×	×
Laleh et al.2013	GDM	Iran	112	3	p	ND)		>0.05	ANCOVA	7			×	×	×	×
Kitajima et al.2001	OGTT +	Japan	146	3	ND	ND			ND	ND	6	ND	ND	ND	ND		ND
Retnakaran et al.2012	non-GDM	Canada	472	3	ND	ND)		ND	ND	7	ND	ND	ND	ND		ND
Ahmad et al. 2006	non-GDM	Malaysia	246	3	ND	ND)		ND	ND	6	ND	ND	ND	ND		ND
					mmol/L	Reference	. 1	LGA	p								
Slagjana et al.2014	non-GDM	Yugoslavia	200	3	$\bar{x}\pm SD$	6.5 ± 1.4 (AGA)	6.0	0±1.0	>0.0<	5 Student t test	5	×	×	×	×	×	×
Son et al.2010	GDM	Korea	104	3	$\mathbf{v} + \mathbf{v}$	5.8±1.1 non-LGA	5.:	5±0.9	0.35	2 Student t test	5	×	×	×	×		×
Hou et al.2014	non-GDM	China	2,790	3		5.30 (AGA) (5.62, 7.10)	•	6.18 19,7.04)	0.01	7 Mann-Whitney U test	7	×	×	×	×		×

The bold font represents statistically significant results.

Abbreviation: Gestational diabetes mellitus(GDM), Positive screenes of Oral Glucose Tolerance test(OGTT+), Confidence interval(CI), Not documented(ND), Multiple logistic regression(MLOR), Analysis of covariance(ANCOVA), Standard deviation (SD), Interquartile range(IQR) and Appropriate for gestational age(AGA).

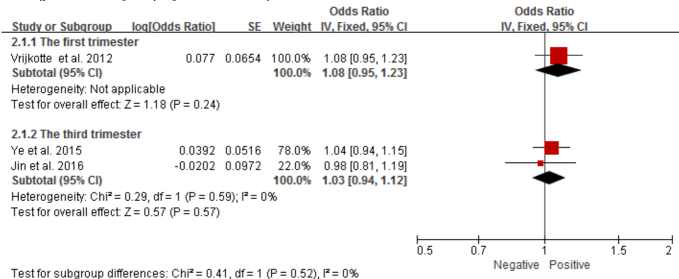
^{*} Result was calculated by comparing the highest quartile with the lowest quartile maternal TC level

[¶] Result was calculated by comparing the highest tertile with the lowest tertile maternal TC level

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels.

Meta-analysis

S8.1 Figure Meta-analysis of adjusted odds ratio for the association between maternal TC levels and LGA



S8.2 Figure Meta-analysis for mean difference of maternal TC levels between LGA and reference groups in the third trimester

	LGA	group		Refere	nce group			Mean Difference	Mea	n Difference	
Study or Subgroup	Mean [mmol/L]	SD [mmol/L]	Total	Mean [mmol/L]	SD [mmol/L]	Total	Weight	IV, Fixed, 95% CI [mmol/L]	IV, Fixed	95% CI [mmol/L]	
Slagjana et al. 2014	6	1	50	6.5	1.4	135	58.0%	-0.50 [-0.86, -0.14]		-	
Son et al. 2010	5.5	0.9	25	5.8	1.1	79	42.0%	-0.30 [-0.73, 0.13]	-		
Total (95% CI) Heterogeneity: Chi ² = 0 Test for overall effect: Z			75			214	100.0%	-0.42 [-0.69, -0.14]	-1 -0.5	0 0.5	1
	•	•							LGA gro	oup Reference group)

High-density lipoprotein cholesterol (HDL-C)

S8.2 Table Results summary of the association of maternal HDL-C levels with LGA

Ctude: ID	Countries	Danulation	Sample	Trimesters	Reported	Effect	Lower	Upper	_	Statistical mathods	Onality	The c	ontrol	of con	ıfoundi	ing fa	ctors
Study ID	Countries	Population	size	Trimesters	measures	size	95%CI	95%CI	р	Statistical methods	Quanty -	a	b	c	d	e	f
Jin et al.2016	China	non-GDM	934	1	ND	ND			ND	ND	7	ND	ND	ND	ND	×	ND
Lei et al.2016	China	General	5,535	2	Crude OR^	0.75	0.63	0.89	ND	Logistic regression	6	×	×	×	×	×	×
Jin et al.2016	China	non-GDM	934	2	ND	ND			ND	ND	7	ND	ND	ND	ND		ND
Di et al.2005	Italy	OGTT+	83	2	ND	ND			ND	ND	5	ND	ND	ND	ND	×	ND
Mossayebi et al.2014	Iran	General	82	3	Crude OR*	0.06	0.01	0.29	ND	Chi-squared test	5	×	×	×	×		×
Retnakaran et al.2012	Canada	non-GDM	472	3	Crude OR	0.89	0.69	1.15	ND	Logistic regression	7	×	×	×	×		×
Ye et al.2015	China	non-GDM	1,204	3	Adjusted OR	0.62	0.47	0.82	ND	MLOR	8				$\sqrt{}$		×
Retnakaran et al.2012	Canada	non-GDM	472	3	Adjusted OR	0.99	0.70	1.39	ND	MLOR	7				$\sqrt{}$		$\sqrt{}$
Jin et al.2016	China	non-GDM	934	3	Adjusted OR	0.79	0.52	1.21	0.281	MLOR	7				×		×
Mossayebi et al.2014	Iran	General	82	3	Adjusted OR*	1.67	0.19	14.29	ND	MLOR	5			×	$\sqrt{}$		$\sqrt{}$
Hou et al.2014*	China	non-GDM	2,790	3	Adjusted OR¶	0.81	0.64	1.04	ND	MLOR	7			×	×		×
Laleh et al.2013	Iran	GDM	112	3	p	ND			>0.05	ANCOVA	7			×	×	×	×
					mmol/L	Reference	i	LGA									
Hou et al.2014	China	non-GDM	2,790	3		1.76 (AGA) (1.52, 2.05)		1.70 8, 1.95)	0.00	Mann-Whitney U test	7	×	×	×	×	V	×
Slagjana et al.2014	Yugoslavia	non-GDM	200	3	$\bar{x} \pm SD$ 1.6	±0.4(non-L0	GA) 1.	.3±0.4	0.00	1 Student t test	5	×	×	×	×	×	×
Son et al.2010	Korea	GDM	104	3	$\bar{x} \pm SD$ 1.7	7 ± 0.5 (non-LC	GA) 1.0	6 ± 0.3	0.23	2 Student t test	5	×	×	×	×		×

The bold font represents statistically significant results.

[^] Results was calculated with self-defined cut-off point: 1.3 mmol/L

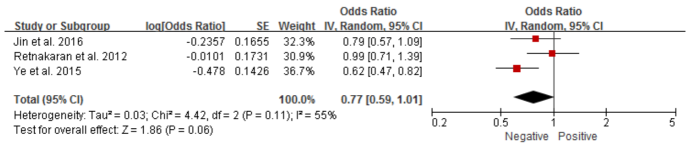
^{*} Result was calculated by comparing the highest quartile with the lowest quartile maternal HDL-C level

[¶] Result was calculated by comparing the highest tertile with the lowest tertile maternal HDL-C level

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels.

Meta-analysis

S8.3 Figure Meta-analysis of adjusted odds ratio for the association between maternal HDL-C levels and LGA in the third trimester



S8.4 Figure Meta-analysis for mean difference of maternal HDL-C levels between LGA and reference groups in the third trimester

	LGA	group		Refere	nce group			Mean Difference		Mean D	ifference		
Study or Subgroup	Mean [mmol/L]	SD [mmol/L]	Total	Mean [mmol/L]	SD [mmol/L]	Total	Weight	IV, Random, 95% CI [mmol/L]	IV	, Random, 9	5% CI [mmol/	<u>L]</u>	
Slagjana et al. 2014	1.3	0.4	50	1.6	0.4	135	53.0%	-0.30 [-0.43, -0.17]		-			
Son et al. 2010	1.6	0.3	25	1.7	0.5	79	47.0%	-0.10 [-0.26, 0.06]		_	 		
Total (95% CI)			75			214	100.0%	-0.21 [-0.40, -0.01]	ı	•			
Heterogeneity: Tau² = 1 Test for overall effect: 2	•	6); I² = 7	'2%					-1 -0).5 LGA group	0 0. Reference o		1	

Sensitivity analysis

S8.5 Figure Sensitivity analysis_ Adjusted odds ratio_ Exclude study adjust for other maternal lipid levels

				Odds Ratio		Od	lds Ratio		
Study or Subgroup	log[Odds Ratio]	SE	Weight	IV, Random, 95% CI		IV, Rar	idom, 95	% CI	
Jin et al. 2016	-0.2357	0.1655	44.0%	0.79 [0.57, 1.09]			-		
Retnakaran et al. 2012	-0.0101	0.1731	0.0%	0.99 [0.71, 1.39]					
Ye et al. 2015	-0.478	0.1426	56.0%	0.62 [0.47, 0.82]		_	•		
Total (95% CI)			100.0%	0.69 [0.54, 0.87]		•	-		
Heterogeneity: Tau ² = 0.0 Test for overall effect: Z =		1 (P = 0.	.27); I² = 1	9%	0.2	0.5	1	2	5
restroi overali ellect. Z=	3.03 (1 - 0.002)					Negati	ve Posit	ive	

Low-density lipoprotein cholesterol (LDL-C)

S8.3 Table Results summary of the association of maternal LDL-C levels with LGA

G. I. ID	G	D 14	Sample,	Trimesters	Reported	Effect	Lower	Upper		Statistical	0 114	The o	ontro	l of co	nfoun	ding f	factors
Study ID	Countries	Population	size	Trimesters	measures	size	95%CI	95%CI	р	methods	Quality	a	b	c	d	e	f
Jin et al.2016	China	non-GDM	934	1	ND	ND			ND	ND	7	ND	ND	ND	ND	×	ND
Jin et al.2016	China	non-GDM	934	2	ND	ND			ND	ND	7	ND	ND	ND	ND		ND
Di et al.2005	Italy	OGTT+	83	2	ND	ND			ND	ND	5	ND	ND	ND	ND	×	ND
Retnakaran et al.2012	Canada	non-GDM	472	3	Crude OR	0.80	0.61	1.05	ND	Logistic regression	1 7	×	×	×	×		×
Mossayebi et al.2014	Iran	General	82	3	Crude OR*	5.80	1.50	22.60	ND	Chi-squared test	5	×	×	×	×		×
Mossayebi et al.2014	Iran	General	77	3	Adjusted OR*	0.80	0.10	4.40	ND	MLOR	5	$\sqrt{}$	$\sqrt{}$	×	$\sqrt{}$		$\sqrt{}$
Hou et al.2014	China	non-GDM	2,790	3	Adjusted OR \P	0.83	0.59	1.17	ND	MLOR	7	$\sqrt{}$	$\sqrt{}$	×	×		×
Ye et al.2015	China	non-GDM	1,204	3	Adjusted OR	1.25	1.06	1.47	ND	MLOR	8	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$		×
Jin et al.2016	China	non-GDM	934	3	Adjusted OR	0.93	0.78	1.11	0.418	3 MLOR	7	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	×		×
Retnakaran et al.2012	Canada	non-GDM	472	3	Adjusted OR	0.98	0.72	1.34	ND	MLOR	7	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$			
Laleh et al.2013	Iran	GDM	112	3	p	ND			>0.05	5 ANCOVA	7		$\sqrt{}$	×	×	×	×
Son et al.2010	Korea	GDM	104	3	ND	ND			ND	ND	5	ND	ND	ND	ND		ND
					mmol/L	Reference	I	LGA									
Hou et al.2014	China	non-GDM	2,790	3		3.07 (AGA) (2.47, 3.74)		2.95 0, 3.65)	0.00	3 Mann-Whitney U test	7	×	×	×	×	$\sqrt{}$	×
Slagjana et al.2014	Yugoslavia	non-GDM	200	3	$\bar{x} \pm SD$	$3.5{\pm}1.2$	3.8	± 1.0	>0.0	5Student t test	5	×	×	×	×	×	×

The bold font represents statistically significant results.

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels.

^{*} Result was calculated by comparing the highest quartile with the lowest quartile maternal LDL-C level

[¶] Result was calculated by comparing the highest tertile with the lowest tertile maternal LDL-C level

Meta-analysis

S8.4 Figure Meta-analysis of adjusted odds ratio for the association between maternal LDL-C levels and LGA in the third trimester

			Odds Ratio		Odds Ratio		
Study or Subgroup	log[Odds Ratio] 9	E Weight	IV, Random, 95% CI	IV,	Random, 95% CI		
Jin et al. 2016	-0.0726 0.089	7 37.3%	0.93 [0.78, 1.11]	_			
Retnakaran et al. 2012	-0.0202 0.157	3 24.1%	0.98 [0.72, 1.33]		-		
Ye et al. 2015	0.2231 0.084	1 38.5%	1.25 [1.06, 1.47]		-	-	
Total (95% CI)		100.0%	1.06 [0.86, 1.30]				
Heterogeneity: Tau² = 0.02 Test for overall effect: Z = (0.05); I² = 8	68%	0.5 0.7 Nec	1 native Positive	1.5	2
				1 gative Positive	1.5	2	

Sensitivity analysis

S8.5 Figure Sensitivity analysis _ Adjusted odds ratio _ The third trimester_ exclude studies adjust for other maternal lipid levels

				Odds Ratio		Odds Ratio		
Study or Subgroup	log[Odds Ratio]	SE	Weight	IV, Random, 95% CI		IV, Random, 95% CI		
Jin et al. 2016	-0.0726	0.0897	49.4%	0.93 [0.78, 1.11]				
Retnakaran et al. 2012	-0.0202	0.1573	0.0%	0.98 [0.72, 1.33]				
Ye et al. 2015	0.2231	0.0841	50.6%	1.25 [1.06, 1.47]			_	
Total (05% CI)			100.0%	1 00 10 01 1 441			_	
Total (95% CI)	4.062 570 46	4 (10 0	100.0%	1.08 [0.81, 1.44]				
Heterogeneity: Tau² = 0.0 Test for overall effect: Z =		1 (P = 0.	02); 1= 8	33%	0.5	0.7 1	1.5	2
restroi everali ellett. 2 –	0.02 (1 = 0.00)					Negative Positive		

Triglycerides (TG)

S8.4 Table Results summary of the association of maternal TG levels with LGA

Study ID	Countries	Population	Sample T	rimesters	Reported	Effect		Upper	P	Statistical methods (Duality	The co	ontrol	of con	foundi	ng fa	ctors
Study ID	Countries	1 opulation	size	i illiestei s	measures	size 9	95%CI	95%CI	1	Statistical methods (Quanty	a	b	c	d	e	f
Jin et al.2016	China	non-GDM	934	1	ND	ND			ND	ND	7	ND	ND	ND	ND	×	ND
Vrijkotte et al.2012	Netherlands	non-GDM	4,008	1	Adjusted OR	1.48	1.23	1.78	ND	MLOR	8		$\sqrt{}$	×	×	×	×
Vrijkotte et al.2012	Netherlands	non-GDM	4,008	1	Crude OR	1.44	1.20	1.71	ND	Logistic regression	8	×	×	×	×	×	×
Lei et al.2016	China	General	5,535	2	Crude OR^	1.60	1.42	2.01	ND	Logistic regression	6	×	×	×	×	×	×
Di et al.2005	Italy	OGTT+	83	2	Crude OR^	5.60	0.93	33.77	ND	Chi-squared test	5	×	×	×	×	×	×
Jin et al.2016	China	non-GDM	934	2	ND	ND			ND	ND	7	ND	ND	ND	ND		ND
Retnakaran et al.2012	Canada	non-GDM	472	3	Crude OR	1.26	0.98	1.62	ND	Logistic regression	7	×	×	×	×		×
Ahmad et al. 2006	Malaysia	non-GDM	246	3	Crude OR^	3.07	1.33	7.08	ND	Chi-squared test	6	×	×	×	×		×
Kitajima et al.2001	Japan	OGTT +	146	3	Crude OR^	14.80	1.59	137.28	0.012	Chi-squared test	6	×	×	×	×		×
Mossayebi et al.2014	Iran	General	154	3	Adjusted OR	1.04	1.02	1.05	ND	MLOR	5		$\sqrt{}$	×	$\sqrt{}$		
Ye et al.2015	China	non-GDM	1,204	3	Adjusted OR	1.15	1.03	1.27	ND	MLOR	8			$\sqrt{}$			×
Retnakaran et al.2012	Canada	non-GDM	472	3	Adjusted OR	0.98	0.70	1.38	ND	MLOR	7	$\sqrt{}$		$\sqrt{}$	$\sqrt{}$		
Jin et al.2016	China	non-GDM	934	3	Adjusted OR	1.13	1.02	1.26	0.025	MLOR	7			$\sqrt{}$	×		×
Hou et al.2014	China	non-GDM	2,790	3	Adjusted OR¶	3.30	1.18	9.27	ND	MLOR	7	$\sqrt{}$		×	×		×
Ahmad et al. 2006	Malaysia	non-GDM	246	3	Adjusted OR^	1.48	1.15	1.93	ND	MLOR	6	×		×			×
Kitajima et al.2001	Japan	OGTT +	146	3	Adjusted OR^	11.60	1.10	122.00	0.040	MLOR	6	×	×	×	×		×
Son et al.2010	Korea	GDM	104	3	Adjusted OR^	4.43	1.33	14.82	ND	MLOR	5				×		×
Schaefer-Graf et al.2008	German	GDM	150	3	p	ND			0.040	MLOR	5		$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	×	×
Laleh et al.2013	Iran	GDM	112	3	p	+			0.040	ANCOVA	7		$\sqrt{}$	×	×	×	×
					mmol/L	Referei	псе	LGA									
Hou et al.2014	China	non-GDM	2,790	3	Median (IQR)	3.02 (AC (2.48, 3.		3.19 (2.61, 3.97)	0.00	0 Mann-Whitney U test	7	×	×	×	×		×
Slagjana et al.2014	Yugoslavia	non-GDM	200	3	$\bar{x}\pm SD$	3.1±1	.1	3.8 ± 1.8	0.012	2 Student t test	5	×	×	×	×	×	×

The bold font represents statistically significant results.

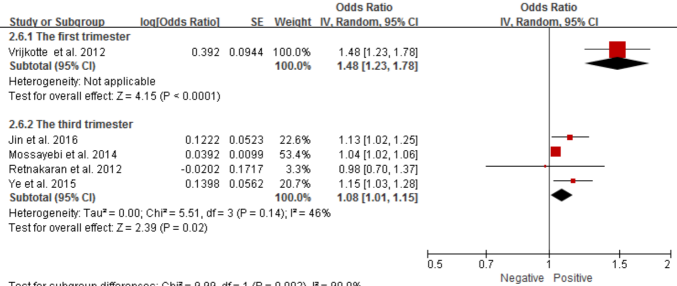
[^] Results was calculated with self-defined cut-off point: Lei et al.2016, 3.49 mmol/L; Di et al.2005, 2.30mmol/L; Ahmad et al. 2006, 2.78mmol/L; Kitajima et al. 2001, 2.92 mmol/L; Son et al. 2010, 3.33mmol/L.

 $[\]P$ Result was calculated by comparing the highest tertile with the lowest tertile maternal TG level

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels.

Meta-analysis

S8.6 Figure Meta-analysis of adjusted odds ratio for the association between maternal TG levels and LGA throughout pregnancy

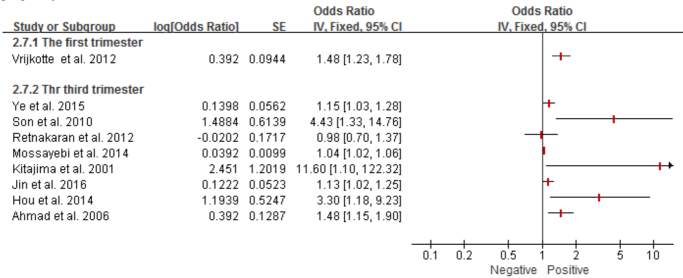


Test for subgroup differences: $Chi^2 = 9.99$, df = 1 (P = 0.002), $I^2 = 90.0\%$

S8.7 Figure Forest plots of crude odds ratio for the association between maternal TG levels and LGA throughout pregnancy

0 1	J	'	Odds Ratio		Odds	Ratio	0 ,
Study or Subgroup	log[Odds Ratio]	SE	IV, Fixed, 95% CI			I, 95% CI	
2.8.1 The first trimeste	er						
Vrijkotte et al. 2012	0.3646	0.093	1.44 [1.20, 1.73]			+	
2.8.2 The second trime	ester						
Di et al. 2005	1.7228	0.916	5.60 [0.93, 33.72]		-	+	\longrightarrow
Lei et al. 2016	0.47	0.0609	1.60 [1.42, 1.80]			+	
2.8.3 The third trimesto	er						
Ahmad et al. 2006	1.1217	0.4268	3.07 [1.33, 7.09]				
Kitajima et al. 2001	2.6946	1.1382	14.80 [1.59, 137.75]				
Retnakaran et al. 2012	0.2311	0.1282	1.26 [0.98, 1.62]			+	
				<u></u>		<u> </u>	
				0.05	0.2 Negative	1 5 Positive	20

S8.8 Figure Forest plots of adjusted odds ratio for the association between maternal TG levels and LGA throughout pregnancy



Sensitivity ananlysis

S8.9 Figure Sensitivity analysis_ Exclude studies adjust for other maternal lipid levels

				Odds Ratio			lds Ratio	
Study or Subgroup	log[Odds Ratio]	SE	Weight	IV, Random, 95% CI		IV, Rai	ndom, 95% CI	
2.6.1 The first trimester							_	
Vrijkotte et al. 2012	0.392	0.0944	100.0%	1.48 [1.23, 1.78]				_
Subtotal (95% CI)			100.0%	1.48 [1.23, 1.78]				
Heterogeneity: Not appli	cable							
Test for overall effect: Z =	= 4.15 (P < 0.0001)							
2.6.2 The third trimeste	г							
Jin et al. 2016	0.1222	0.0523	53.6%	1.13 [1.02, 1.25]				
Mossayebi et al. 2014	0.0392	0.0099	0.0%	1.04 [1.02, 1.06]				
Retnakaran et al. 2012	-0.0202	0.1717	0.0%	0.98 [0.70, 1.37]				
Ye et al. 2015	0.1398	0.0562	46.4%	1.15 [1.03, 1.28]			——	
Subtotal (95% CI)			100.0%	1.14 [1.06, 1.23]			•	
Heterogeneity: Tau ² = 0.1	00; Chi² = 0.05, df =	1 (P = 0.	82); $I^2 = 0$	%				
Test for overall effect: Z=	3.41 (P = 0.0007)	•						
	,,							
					<u></u>	- 	1 15	
					0.5	0.7	1 1.5	2
Toot for outpareup differe		4 (D -	0.043 12	- 04 000		Negat	ive Positive	

Test for subgroup differences: $Chi^2 = 6.60$, df = 1 (P = 0.01), $I^2 = 84.8\%$

Free fatty acids (FFAs)

S8.5 Table Results summary of the association of maternal FFAs levels with LGA

C4 J ID	C	D	Sample	T	Reported	T-664 -:		Statistical	01'4	The o	contro	ol of c	onfou	nding f	actors	, TT24
Study ID	Countries	Population	size	Trimesters	measures	Effect size	р	methods	Quality	a	b	c	d	e	f	- Unit
Schaefer-Graf et al.2008	German	GDM	150	3	p	ND	0.008	MLOR	5				$\sqrt{}$	×	×	μmol/L
Kitajima et al.2001	Japan	OGTT +	146	3	ND	ND	ND	ND	6	×	×	×	×	$\sqrt{}$	×	ND



Supplementary 9 Data analysis for Small for gestational age (SGA)

Total cholesterol (TC)

S9.1 Table Results summary of the association of maternal TC levels with SGA

Study ID	Countries	Danulation	Sample	Trimesters	Reported	Effect	Lower	Upper	n	Statistical	Quality	The c	ontro	of co	nfound	ling fa	actors
Study ID	Countries	Population	size	Timesters	measures	size	95%CI	95%CI	Р	methods	Quanty	a	b	c	d	e	f
Vrijkotte et al.2012	Netherlands	non-GDM	4,008	1	Crude OR	0.97	0.85	1.10	ND	Logistic regression	8	×	×	×	×	×	×
Vrijkotte et al.2012	Netherlands	non-GDM	4,008	1	Adjusted OR	0.98	0.86	1.12	ND	MLOR	8		$\sqrt{}$	×	×	×	×
Jin et al.2016	China	non-GDM	934	1	ND	ND			ND	ND	7	ND	ND	ND	ND	$\sqrt{}$	ND
Jin et al.2016	China	non-GDM	934	2	ND	ND			ND	ND	7	ND	ND	ND	ND	$\sqrt{}$	ND
Ye et al.2015	China	non-GDM	912	3	Adjusted OR	0.94	0.74	1.20	ND	MLOR	8				$\sqrt{}$		×
Jin et al.2016	China	non-GDM	934	3	Adjusted OR	1.12	0.80	1.56	0.520	MLOR	7		$\sqrt{}$	$\sqrt{}$	×	$\sqrt{}$	×
Slagjana et al.2014	Yugoslavia	non-GDM	200	3	p			:	>0.05	Student t test	5	×	×	×	×	×	×

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels. Abbreviation: Gestational diabetes mellitus(GDM), Confidence interval(CI), Not documented(ND), Multiple logistic regression(MLOR).

S9.1 Figure Meta-analysis of adjusted odds ratio for the association between maternal TC levels and SGA throughout pregnancy

				Odds Ratio		Odds	Ratio		
Study or Subgroup	log[Odds Ratio]	SE	Weight	IV, Fixed, 95% CI		IV, Fixed	1, 95% CI		
4.1.1 The first trimes	ster						L		
Vrijkotte et al. 2012	-0.0202	0.0666	100.0%	0.98 [0.86, 1.12]		_			
Subtotal (95% CI)			100.0%	0.98 [0.86, 1.12]		-	-		
Heterogeneity: Not ap	plicable								
Test for overall effect:	Z = 0.30 (P = 0.76)	l							
4.1.2 The third trimes	ster								
Jin et al. 2016	-0.0619	0.1221	66.4%	0.94 [0.74, 1.19]			 		
Ye et al. 2015	0.1133	0.1717	33.6%	1.12 [0.80, 1.57]			-		
Subtotal (95% CI)			100.0%	1.00 [0.82, 1.21]					
Heterogeneity: Chi ² =	0.69, df = 1 (P = 0.4	41); $I^2 = 0$)%						
Test for overall effect:	Z = 0.03 (P = 0.98)								
					——	+			—
					0.5	0.7	1	1.5	2
Took for our barrows die	favorana ObiZ — O C	10 de 4	(D = 0.00)	17 - 00/		Negative	Positive		

Test for subgroup differences: Chi² = 0.02, df = 1 (P = 0.89), I^2 = 0%

High-density lipoprotein cholesterol (HDL-C)

S9.2 Table Results summary of the association of maternal HDL-C levels with SGA

Ct., J., ID	Comtrios	Danulation	Sample	Trimesters	Reported	Effect	Lower	Upper	_	Statistical	O 1:4	The c	ontro	l of co	onfour	nding f	factors
Study ID	Countries	Population	size	Trimesters	measures	size	95%CI	95%CI	Р	methods	Quality	a	b	c	d	e	f
Jin et al.2016	China	non-GDM	934	1	Adjusted OR	1.4	0.32	5.38	ND	MLOR	7	$\sqrt{}$			×	$\sqrt{}$	×
Lei et al.2016	China	General	5,535	2	Crude OR^	1.13	3 0.80	1.61	ND	Logistic regression	6	×	×	×	×	×	×
Jin et al.2016	China	non-GDM	934	2	Adjusted OR	1.88	8 0.47	7.59	ND	MLOR	7	$\sqrt{}$		$\sqrt{}$	×	$\sqrt{}$	×
Ye et al.2015	China	non-GDM	912	3	Adjusted OR	1.5	7 0.87	2.83	ND	MLOR	8	$\sqrt{}$			$\sqrt{}$	$\sqrt{}$	×
Jin et al.2016	China	non-GDM	934	3	Adjusted OR	3.1	5 1.15	8.65	0.02	6 MLOR	7	$\sqrt{}$		$\sqrt{}$	×	$\sqrt{}$	×
Slagjana et al.2014	Yugoslavia	non-GDM	200	3	p				>0.0	5 Student t test	5	×	×	×	×	×	×

The bold font represents statistically significant results.

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels.

Abbreviation: Gestational diabetes mellitus(GDM), Confidence interval(CI), Not documented(ND), Multiple logistic regression(MLOR).

S9.2 Figure Meta-analysis of adjusted odds ratio for the association between maternal HDL-C levels and SGA throughout pregnancy

				Odds Ratio			Odds	Ratio		
Study or Subgroup	log[Odds Ratio]	SE	Weight	IV, Random, 95% CI			IV, Rando	om, 95% CI		
4.2.1 The first trimes	ster							l <u> </u>		
Jin et al. 2016	0.3436	0.7567	100.0%	1.41 [0.32, 6.21]						-
Subtotal (95% CI)			100.0%	1.41 [0.32, 6.21]						
Heterogeneity: Not ap	plicable									
Test for overall effect:	Z = 0.45 (P = 0.65)	ı								
4.2.2 The second trir	nester							_		
Jin et al. 2016	0.6313	0.7073	100.0%	1.88 [0.47, 7.52]						_
Subtotal (95% CI)			100.0%	1.88 [0.47, 7.52]						
Heterogeneity: Not ap	oplicable									
Test for overall effect:	Z= 0.89 (P = 0.37)	1								
4.2.3 The third trimes	ster									
Jin et al. 2016	1.1474	0.5141	32.1%	3.15 [1.15, 8.63]						
Ye et al. 2015	0.4511	0.3012	67.9%	1.57 [0.87, 2.83]			-			
Subtotal (95% CI)			100.0%	1.96 [1.04, 3.71]					-	
Heterogeneity: Tau ² =	0.06; Chi ² = 1.37,	df= 1 (P :	= 0.24); l ²	= 27%						
Test for overall effect:	Z = 2.08 (P = 0.04)									
					<u></u>			<u> </u>	<u> </u>	
					0.1	0.2	0.5	1 2	5	10
Toot for cubarous diff	foroncoe: Chi z – 0 1	16 df = 2	/D = 0.02	\ IZ = 0.06			Negative	Positive		

Test for subgroup differences: $Chi^2 = 0.16$, df = 2 (P = 0.92), $I^2 = 0\%$

[^] Results was calculated with self-defined cut-off point: 1.3 mmol/L

Low-density lipoprotein cholesterol (LDL-C)

S9.3 Table Results summary of the association of maternal LDL-C levels with SGA

C4 J ID	Communica	Danulation	Sample	T	Reported	Effect	Lower		Statistical	Oa1!4	The c	ontrol	of con	foundi	ng fa	ctors
Study ID	Countries	Population	size	Trimesters	measures	size	95%CI	95%CI ^p	methods	Quality	a	b	c	d	e	f
Jin et al.2016	China	non-GDM	934	1	ND	ND		ND	ND	7	ND	ND	ND	ND	$\sqrt{}$	ND
Jin et al.2016	China	non-GDM	934	2	ND	ND		ND	ND	7	ND	ND	ND	ND	$\sqrt{}$	ND
Ye et al.2015	China	non-GDM	912	3	Adjusted OR	0.75	0.50	1.14 ND	MLOR	8	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$			×
Jin et al.2016	China	non-GDM	934	3	Adjusted OR	1.16	0.71	1.89 0.565	MLOR	7	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	×	$\sqrt{}$	×
Slagjana et al.2014	Yugoslavia	non-GDM	200	3	p			>0.05	Student t test	5	×	×	×	×	×	×

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels. Abbreviation: Gestational diabetes mellitus(GDM), Confidence interval(CI), Not documented(ND), Multiple logistic regression(MLOR).

S9.3 Figure Meta-analysis of adjusted odds ratio for the association between maternal LDL-C levels and SGA in the third trimester

Study or Subgroup log[Odds Ratio] SE Weight IV, Random, 95% CI IV, Random, 95% CI Jin et al. 2016 0.1484 0.2505 44.8% 1.16 [0.71, 1.90] 1.16 [0.71, 1.90] Ye et al. 2015 -0.2877 0.2069 55.2% 0.75 [0.50, 1.13] 0.75 [0.50, 1.13] Total (95% CI) 100.0% 0.91 [0.60, 1.39] 0.1 0.2 0.5 1 2 5 10 Heterogeneity: Tau² = 0.04; Chi² = 1.80, df = 1 (P = 0.18); I² = 44% 0.1 0.2 0.5 1 2 5 10 Test for overall effect: Z = 0.43 (P = 0.67) Negative Positive					Odds Ratio		Odds Ratio		
Ye et al. 2015 -0.2877 0.2069 55.2% 0.75 [0.50, 1.13] Total (95% CI)	Study or Subgroup	log[Odds Ratio]	SE	Weight	IV, Random, 95% CI		IV, Random, 95% CI		
Total (95% CI) Heterogeneity: Tau² = 0.04; Chi² = 1.80, df = 1 (P = 0.18); I² = 44% Test for overall effect: 7 = 0.43 (P = 0.67) Test for overall effect: 7 = 0.43 (P = 0.67)	Jin et al. 2016	0.1484	0.2505	44.8%	1.16 [0.71, 1.90]		_		
Heterogeneity: Tau ² = 0.04; Chi ² = 1.80, df = 1 (P = 0.18); I ² = 44% Test for overall effect: 7 = 0.43 (P = 0.67) Test for overall effect: 7 = 0.43 (P = 0.67)	Ye et al. 2015	-0.2877	0.2069	55.2%	0.75 [0.50, 1.13]				
Test for overall effect: 7 = 0.43 (P = 0.67) 0.1 0.2 0.5 1 2 5 10	Total (95% CI)			100.0%	0.91 [0.60, 1.39]		-		
Test for overall effect: $7 = 0.43$ ($P = 0.67$)				= 0.18); l²	2 = 44%	0.1 0.2	0.5 1 2	 5	— 10
	Test for overall effect:	Z = 0.43 (P = 0.67)					Negative Positive	-	

Triglycerides (TG)

S9.4 Table Results summary of the association of maternal TG levels with SGA

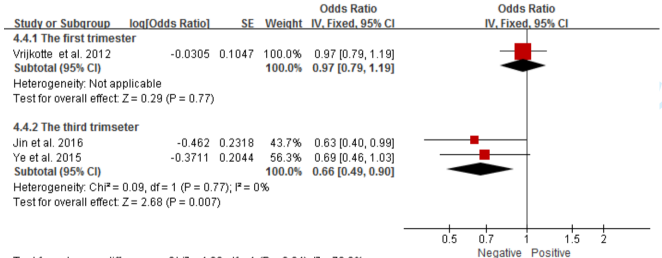
C4 J ID	Carratria	Danulation	Sample	Trimesters	Reported	Effect	Lower	Upper		Statistical	Oa1:4	The	contro	ol of co	nfound	ing fa	ctors
Study ID	Countries	Population	size	Trimesters	measures	size	95%CI	95%CI	þ	methods	Quality	a	b	c	d	e	f
Jin et al.2016	China	non-GDM	934	1	ND	ND)		ND	ND	7	ND	ND	ND	ND	$\sqrt{}$	ND
Vrijkotte et al.2012	Netherlands	non-GDM	4,008	1	Crude OR	1.06	0.87	1.29	ND	Logistic regression	8	×	×	×	×	×	×
Vrijkotte et al.2012	Netherlands	non-GDM	4,008	1	Adjusted OR	0.97	0.79	1.19	ND	MLOR	8			×	×	×	×
Jin et al.2016	China	non-GDM	934	2	ND	ND	•		ND	ND	7	ND	ND	ND	ND	$\sqrt{}$	ND
Lei et al.2016	China	General	5,535	2	Crude OR^	1.51	1.08	2.12	ND	Logistic regression	6	×	×	×	×	×	×
Ye et al.2015	China	non-GDM	912	3	Adjusted OR	0.69	0.47	1.03	ND	MLOR	8			$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	×
Jin et al.2016	China	non-GDM	934	. 3	Adjusted OR	0.63	0.40	0.99	0.046	MLOR	7				×		×
Slagjana et al.2014	Yugoslavia	non-GDM	200	3	p				0.012	Student t test	5	×	×	×	×	×	×
TTI 1 11 C	11	. 1.															

The bold font represents statistically significant results.

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels.

Abbreviation: Gestational diabetes mellitus(GDM), Confidence interval(CI), Not documented(ND), Multiple logistic regression(MLOR).

S9.4 Figure Meta-analysis of adjusted odds ratio for the association between maternal TG levels and SGA throughout pregnancy



Test for subgroup differences: Chi² = 4.20, df = 1 (P = 0.04), I^2 = 76.2%

[^] Results was calculated with self-defined cut-off point: 3.49 mmol/L

Supplementary 10 Data analysis for Macrosomia

Total cholesterol (TC)

S10.1 Table Results summary of the association of maternal TC levels with macrosomia

Ct.,.J., ID	Countries	Donulation	Sample	т:	Reported	Effect 1	Lower Up	per		Statistical methods	Onality	The	con	trol (of co	nfou	nding	g fac	tors
Study ID	Countries	Population	size	1 11.	measures	size 9	95%CI95	%CI	þ	Staustical methods	Quality	a	b	c	d	e	f	g	h
Jin et al.2016	China	non-GDM	934	1	ND	ND			ND	ND	7	ND	ND	ND	ND	ND	ND	×	ND
Clausen et al.2005	Norway	General	1,037	2	Crude OR*	1.10	0.60	2.00	ND	Logistic regression	8	×	×	×	×	×	×		×
Clausen et al.2005	Norway	General	1,037	2	Adjusted OR*	1.10	0.60	2.00	ND	MLOR	8	×	×		×	×	×		×
Zhou et al.2012	China	General	1,000	2	P				>0.05	Non-parametric Mann-Whitney Test	5	×	×	×	×	×	×	×	×
Jin et al.2016	China	non-GDM	934	2	ND	ND			ND	ND	7	ND	ND	ND	ND	ND	ND		ND
Jin et al.2016	China	non-GDM	934	3	Adjusted OR	0.99	0.81	1.21	0.903	MLOR	7	×					×		×
Laleh et al.2013	Iran	GDM	112	3	P	ND			>0.05	Bonferroni multiple comparison test	7	×	×			×	×	×	×
Mossayebi et al.2014	Iran	General	154	3	ND	ND			ND	ND	5	ND	ND	ND	ND	ND	ND		ND

^{*} Result was calculated by comparing the highest quartile with the lowest quartile maternal TC level

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels.

High-density lipoprotein cholesterol (HDL-C)

S10.2 Table Results summary of the association of maternal HDL-C levels with macrosomia

C. I. ID	a	D 14	Sample	m •	Reported	Effect	Lower U	pper			0 114		con	trol o	of cor	nfou	nding	g fac	tors
Study ID	Countries	Population	size	Tri.	measures	size	95%CI95	%CI	p	Statistical methods	Quality	a	b	c	d	e	f	g	h
Jin et al.2016	China	non-GDM	934	1	Adjusted OR	0.51	0.19	1.36	0.178	MLOR	7	×					×		×
Zawiejska et al. 2008	Poland	GDM	357	2	Crude RR	0.59	0.32	1.02	ND	Chi-squared test	5	×	×	×	×	×	×	×	×
Clausen et al.2005	Norway	General	1,025	2	Crude OR*	0.30	0.20	0.60	ND	Logistic regression	8	×	×	×	×	×	×		×
Clausen et al.2005	Norway	General	1,025	2	Adjusted OR*	0.30	0.20	0.60	ND	MLOR	8	×	×		×	×	×	$\sqrt{}$	×
Zhou et al.2012	China	General	1,000	2	Adjusted OR^	0.61	0.38	0.98	ND	MLOR	5	×	×	$\sqrt{}$			×	×	×
Jin et al.2016	China	non-GDM	934	2	Adjusted OR	0.25	0.09	0.73	0.011	MLOR	7	×	$\sqrt{}$				×	$\sqrt{}$	×
Jin et al.2016	China	non-GDM	934	3	Adjusted OR	0.46	0.22	0.94	0.034	MLOR	7	×					×		×
Laleh et al.2013	Iran	GDM	112	3	p	ND			>0.05	Bonferroni multiple comparison test	7	×	×	$\sqrt{}$		×	×	×	×
Mossayebi et al.2014	Iran	General	154	3	ND	ND			ND	ND	5	ND	ND	ND	ND	ND	ND	V	ND

The bold font represents statistically significant results.

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels.

S10.1 Figure Forest plots of adjusted odds ratio for the association between maternal HDL-C levels and macrosomia throughout pregnancy

			Odds Ratio		Odds	Ratio		
Study or Subgroup	log[Odds Ratio]	SE	IV, Random, 95% CI		IV, Rando	om, 95% CI		
3.2.1 The first trimes	ter							
Jin et al. 2016	-0.6733	0.5004	0.51 [0.19, 1.36]			_		
3.2.2 The second trin	nester							
Couch et al.1998	-1.204	0.2069	0.30 [0.20, 0.45]					
Jin et al. 2016	-1.3863	0.5213	0.25 [0.09, 0.69]	•				
Zhou et al. 2012	-0.4943	0.2419	0.61 [0.38, 0.98]			-		
3.2.3 The third trimes	etor							
Jin et al. 2016	-0.7765	0.2646	0.46 [0.23, 0.94]			.		
Jili et al. 2010	-0.7703	0.3040	0.40 [0.25, 0.54]					
				—		 		—
				0.1	0.2 0.5	1 2	5	10
					Negative	Positive		

[^] Results was calculated with self-defined cut-off point: 2.205mmol/L

^{*} Result was calculated by comparing the highest quartile with the lowest quartile maternal HDL-C level

Low-density lipoprotein cholesterol (LDL-C)

S10.3 Table Results summary of the association of maternal LDL-C levels with macrosomia

Study ID	Countries	Population	Sample Tri.	Reported	Effect I	Lower Up	per	n	Statistical methods	Quality	The	con	trol (of co	nfoui	nding	g fac	tors
Study 1D	Countries	1 opulation	size 111.	measures	size 9	95%CI95%	%CI	Р	Statistical methods	Quanty	a	b	c	d	e	f	g	h
Jin et al.2016	China	non-GDM	934 1	ND	ND			ND	ND	7	ND	ND	ND	ND	ND	ND	×	ND
Clausen et al.2005	Norway	General	1,018 2	Crude OR*	2.20	1.20	4.00	ND	Logistic regression	8	×	×	×	×	×	×		×
Clausen et al.2005	Norway	General	1,018 2	Adjusted OR*	2.10	1.20	3.90	ND	MLOR	8	×	×		×	×	×		×
Zhou et al.2012	China	General	1,000 2	p			:	>0.05	Non-parametric Mann-Whitney Test	5	×	×	×	×	×	×	×	×
Jin et al.2016	China	non-GDM	934 2	ND	ND			ND	ND	7	ND	ND	ND	ND	ND	ND		ND
Jin et al.2016	China	non-GDM	934 3	Adjusted OR	0.93	0.69	1.25	0.621	MLOR	7	×					X		×
Laleh et al.2013	Iran	GDM	112 3	p	ND			>0.05	Bonferroni multiple comparison test	7	×	×			×	×	×	×
Mossayebi et al.2014	Iran	General	154 3	ND	ND			ND	ND	5	ND	ND	ND	ND	ND	ND		ND

The bold font represents statistically significant results.

Abbreviation: Gestational diabetes mellitus(GDM), Positive screenes of Oral Glucose Tolerance test(OGTT+), Confidence interval(CI), Not documented(ND), Multiple logistic regression(MLOR), Analysis of covariance(ANCOVA), Standard deviation (SD), Interquartile range(IQR) and Appropriate for gestational age(AGA).

Review

^{*} Result was calculated by comparing the highest quartile with the lowest quartile maternal LDL-C level

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels.

Triglycerides (TG)

S10.4 Table Results summary of the association of maternal TG levels with macrosomia

Study ID	Countries	Population	Sample Tri	Reported		Lower U _l		n	Statistical methods	Quality	The	e con	trol (of co	nfou	ndin	g fac	tors
Study ID	Countries	1 opulation	size	measures	size 9	5%CI95	%CI	Р	Statistical methods	Quanty	a	b	c	d	e	f	g	h
Jin et al.2016	China	non-GDM	934 1	ND	ND			ND	ND	7	ND	ND	ND	ND	ND	ND	×	ND
Clausen et al.2005	Norway	General	988 2	Crude OR*	2.90	1.40	5.90	ND	Logistic regression	8	×	×	×	×	×	×		×
Clausen et al.2005	Norway	General	988 2	Adjusted OR*	2.90	1.40	5.90	ND	MLOR	8	×	×		×	×	×		×
Zhou et al.2012	China	General	1,000 2	p				>0.05	Non-parametric Mann-Whitney Test	5	×	×	×	×	×	×	×	×
Jin et al.2016	China	non-GDM	934 2	ND	ND			ND	ND	7	ND	ND	ND	ND	ND	ND		ND
Mossayebi et al.2014	Iran	General	154 3	Adjusted OR	1.04	1.02	1.07	ND	MLOR	5	×	×			×			$\sqrt{}$
Jin et al.2016	China	non-GDM	934 3	Adjusted OR	1.19	1.02	1.39	0.024	MLOR	7	×					×		×
Lin et al.2013	China	General	ND NI	OR^	2.20	1.54	3.14	ND	ND	NA	ND	ND	ND	ND	ND	ND	ND	ND
Laleh et al.2013	Iran	GDM	112 3	p	+			0.001	Bonferroni multiple comparison test	7	×	×			×	×	×	×

The bold font represents statistically significant results.

Confounding factors: a. Maternal age; b. Pre-pregnancy BMI; c. Gestational weight gain; d. Maternal glucose level; e. pre-term birth; f. Maternal lipid levels.

S10.2 Figure Meta-analysis of adjusted odds ratio for the association between maternal TG levels and macrosomia

				Odds Ratio		Odds	Ratio		
Study or Subgroup	log[Odds Ratio]	SE	Weight	IV, Random, 95% CI		IV, Rando	om, 95% CI		
Jin et al. 2016	0.174	0.0786	33.3%	1.19 [1.02, 1.39]				-	
Mossayebi et al. 2014	0.0392	0.0099	66.7%	1.04 [1.02, 1.06]					
Total (95% CI)			100.0%	1.09 [0.96, 1.23]			~		
Heterogeneity: Tau² = 0 Test for overall effect: Z		=1 (P=0).09); l²=	65%	0.5	0.7 Negative	1 Positive	1.5	<u></u>

[^] Results was calculated with self-defined cut-off point: 2.27 mmol/L

^{*} Result was calculated by comparing the highest quartile with the lowest quartile maternal TG level

S10.3 Figure Forest plots of adjusted odds ratio for the association between maternal TG levels and macrosomia

			Odds Ratio	Odds Ratio
Study or Subgroup	log[Odds Ratio]	SE	IV, Fixed, 95% CI	IV, Fixed, 95% CI
3.1.1 The second trimester				
Clausen et al. 2005	1.0647	0.3716	2.90 [1.40, 6.01]	
3.1.2 The third trimester				
Jin et al. 2016			1.19 [1.02, 1.39]	
Mossayebi et al. 2014	0.0392	0.0099	1.04 [1.02, 1.06]	•
3.1.3 Unkown trimester				
Lin et al. 2013	0.7885	0.182	2.20 [1.54, 3.14]	
				0.1 0.2 0.5 1 2 5 10
				Negative Positive