**Mismatch Between Poor Fetal Growth and Rapid Postnatal Weight Gain in the First 2 Years of Life Is Associated with Higher Blood Pressure and Insulin Resistance without Increased Adiposity in Childhood: The GUSTO Cohort Study**

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# Abstract

**Background:** Using longitudinal ultrasounds as an improved fetal growth marker, we aimed to investigate if fetal growth deceleration followed by rapid postnatal weight gain is associated with childhood cardiometabolic risk biomarkers in a contemporary well-nourished population.

**Methods:** We defined fetal growth deceleration (FGD) as ultrasound-measured 2nd-3rd-trimester abdominal circumference decrease by ≥0.67 standard deviation score [SDS] and rapid postnatal weight gain (RPWG) as 0-2-year-old weight increase by ≥0.67 SDS. In the GUSTO mother-offspring cohort, we grouped 797 children into 4 groups of FGD-only (14.2%), RPWG-only (23.3%), both (mismatch, 10.7%) or neither (reference, 51.8%). Adjusting for confounders and comparing to the reference group, we tested associations of these growth groups with childhood cardiometabolic biomarkers: MRI-measured abdominal fat (n=262), liver fat (n=216), intramyocellular lipids (n=227), Quantitative Magnetic Resonance-measured overall body fat % (BF%) (n=310), homeostasis model assessment of insulin resistance (HOMA-IR) (n=323), arterial wall thickness (n=422) and stiffness (n=443), and blood pressure trajectories (ages 3-6y).

**Results:** Mean±SD birthweights were: FGD-only (3.11±0.38kg), RPWG-only (3.03±0.37kg), mismatch (2.87±0.31kg), reference (3.30±0.36kg). FGD-only children had elevated blood pressure trajectories without correspondingly increased BF%. RPWG-only children had altered body fat partitioning, higher BF% (B=4.26% 95%CI [2.34,6.19]), HOMA-IR (0.28units [0.11,0.45]), and elevated blood pressure trajectories. Mismatch children did not have increased adiposity, but had elevated ectopic fat, elevated HOMA-IR (0.29units [0.04,0.55]), and the highest blood pressure trajectories. Associations remained even after excluding small-for-gestational-age infants from analyses.

**Conclusions:** Fetal growth deceleration coupled with rapid postnatal weight gain was associated with elevated childhood cardiometabolic risk biomarkers without correspondingly increased BF%.

**Key words:** cohort study, fetal growth, postnatal growth, mismatch, cardiometabolic risk, adiposity, insulin resistance, body composition, cardiovascular, blood pressure

# Key messages

* We used dynamic growth information from longitudinal fetal ultrasounds rather than birthweight-for-gestational-age as a marker of poor fetal growth.
* Children who experienced 2nd–3rd trimester fetal growth deceleration had elevated blood pressure trajectories without a corresponding increase in body fat %, arterial thickness, or arterial stiffness, suggesting that poor fetal growth might affect blood pressure not through adiposity or changes in arterial structure, but possibly through poorer nephrogenesis.
* Children who experienced 0-2-year-old rapid postnatal weight gain, regardless of their fetal growth rates, had elevated abdominal fats, intramyocellular lipids, insulin resistance, and blood pressure trajectories in early childhood (ages 3-6y), which could lead to greater cardiometabolic risk if these biomarkers track to adulthood.
* Children who experienced a mismatch of fetal growth deceleration followed by rapid postnatal weight gain had the highest blood pressure trajectories and multiple elevated cardiometabolic risk biomarkers without a corresponding increase in overall body fat %, suggesting that rapid postnatal weight gain might act as a “second-hit” and increase cardiometabolic risk in children who had experienced poor fetal growth.
* Our findings highlight the importance of monitoring longitudinal growth patterns right from the fetal stage.

#  Introduction

The first 1000 days of life is a developmentally plastic and nutritionally sensitive period which may program future cardiometabolic risk (1). From an evolutionary perspective, poor nutritional cues in-utero enable predictive adaptive responses to maximize chances of survival in the predicted poor postnatal nutritional environment. However, when the postnatal nutritional environment is richer than predicted, predictive adaptive responses may become maladaptive and lead to increased cardiometabolic risk (2–4). This phenomenon is known as “developmental mismatch”, which has been supported by animal studies investigating mismatch between maternal and postnatal nutrition as well as epidemiological studies investigating mismatch between fetal and postnatal growth (2–10).

Poor fetal growth, rapid postnatal weight gain (RPWG), and a combination of these two growth patterns have been associated with increased cardiometabolic risk (5,8,9,11–14). However, earlier studies often used low birthweight or small-for-gestational-age (SGA) as a crude proxy of poor fetal growth (5,8). These definitions might incorrectly classify neonates who were constitutionally small as having poor fetal growth (15). Hence, we studied fetal growth deceleration (FGD), defined as downward crossing of at least one major centile band (≥0.67 standard deviation score [SDS]) in fetal abdominal circumference charts between the 2nd and 3rd trimester. Crossing down by at least one major centile band (0.67 SDS units) has been commonly used to assess growth deceleration while crossing up by at least one major centile band has been used to assess growth acceleration, in both fetal and postnatal life (13,16). In addition, earlier studies on developmental mismatch has focused on extreme nutritional stresses in-utero such as famine, intrauterine ligation, or extreme caloric restriction in animal models (6,7,9). It is less clear if the adverse cardiometabolic risk associated with developmental mismatch manifest even in contemporary, well-nourished populations.

We aimed to evaluate the associations of FGD alone, RPWG alone, and a combination of these growth patterns with early cardiometabolic alterations in childhood in an Asian mother-offspring cohort. We used an extensive panel of childhood cardiometabolic biomarkers: adiposity outcomes (anthropometry, BF%, abdominal subcutaneous adipose tissue, visceral adipose tissue, intramyocellular lipids, liver fat), metabolic outcomes (fasting plasma glucose [FPG], fasting plasma insulin, insulin resistance), and cardiovascular outcomes (blood pressure, arterial thickness and stiffness). More sophisticated markers such as BF% and body fat partitioning to the abdomen, muscles, and liver have been shown to be strong, independent correlates of cardiometabolic risk, above and beyond that of BMI alone (17–20). We hypothesize that the mismatch group with FGD followed by RPWG would be associated with the greatest alterations in cardiometabolic risk biomarkers, even in early childhood.

# Materials and Methods

***Study population***

Children were part of the Growing Up in Singapore Towards healthy Outcomes (GUSTO) longitudinal mother-offspring cohort, detailed previously (21). Pregnant women in the first trimester were recruited (June 2009-September 2010) from KK Women’s and Children’s Hospital and National University Hospital, Singapore. Of 3751 women, 2034 were eligible, 1247 were recruited (response rate 61.3%), and 797 singletons with both fetal and postnatal growth data available were included (Supplementary Fig.S1). Preterm (<37 completed weeks gestation) infants were excluded as World Health Organization (WHO) growth charts used to derive weight z-scores were not suitable for use in preterm infants. Data for different cardiometabolic assessments were not available for all children due to differential consent for measurement, data quality control, and loss to follow-up. Precise sample sizes for different cardiometabolic outcomes are reported (Supplementary Fig.S1). Written informed consent was obtained. The National Healthcare Group Domain Specific Review Board and SingHealth Centralized Institutional Review Board approved the study.

***Exposures***

Fetal abdominal circumference, a good indicator of fetal growth (22), was measured by trained ultrasonographers in the second (mean gestational age (±SD): 20.3±0.8wks) and third (32.9±0.7wks) trimesters. SDS were derived using the INTERGROWTH-21st standards (23). FGD was defined as a downward crossing of fetal abdominal circumference by at least one major centile band (change in SDS by ≥0.67 units) from the second to third trimester scan, similar to previous studies (24–26). Crossing down by at least one major centile band (0.67 SDS units) has been commonly used to assess growth deceleration while crossing up by at least one major centile band has been used to assess growth acceleration, in both fetal and postnatal life (13,16).

Birthweights were extracted from hospital medical records and weights (in g) at age 2 years were measured (SECA803 Weighing Scale) (27). SDS were derived according to WHO child growth standards (28), using the LMS method (29). RPWG was defined as an increase in weight by at least one major centile band (change in SDS by ≥0.67 units) from birth to age 2 years (13).

Four growth groups were defined, depending on whether the child experienced FGD-only (14.2%), RPWG-only (23.3%), both FGD and RPWG (mismatch group) (10.7%), or neither (reference group) (51.8%).

***Cardiometabolic Risk Biomarkers***

At ages 4.5 and 6 years, weight (SECA803 Weighing Scale) and standing height (SECA213 stadiometer) were measured in 674 children (27). SDS of weight, height, and BMI were derived using WHO growth standards (28). At age 6 years, high precision and accurate overall fat mass were obtained by Quantitative Magnetic Resonance (QMR) (EchoMRI-Adolescent Humans Body Composition Analyzer, EchoMRI Corporation, Singapore) in a subset of 310 children (30). BF% was calculated by: [fat mass (kg) / total body weight (kg)]\*100%

At age 4.5 years, abdominal MRI was performed without sedation using a Siemens Skyra 3T magnetic resonance scanner to derive volumes of abdominal subcutaneous adipose tissue and visceral adipose tissue in cubic centimeters (cc) in 262 children (31,32). Intramyocellular lipids and liver fat were assessed by proton magnetic resonance spectroscopy (1H-MRS) in 227 and 216 children respectively (33). Intramyocellular lipids was expressed as a percentage of the water signal and is proportional to lipid accumulation within skeletal muscles. Liver fat percentage by weight was calculated by averaging across the right and left liver lobe scans (Detailed MRI and MRS protocols in supplementary section).

At age 6 years, after an overnight fast, venous fasting plasma glucose (FPG) was measured using the hexokinase enzymatic method (Abbott Architect c8000 analyzer at KK Women’s and Children’s Hospital and Beckman AU5800 analyzer at National University Hospital) in 429 children. Fasting insulin was measured using a sandwich immunoassay (Beckman DXL800 analyzer, Beckman Coulter) in 324 children. In 323 children, Homeostasis model assessment (HOMA) of insulin resistance (HOMA-IR) was calculated as follows: HOMA-IR = [fasting insulin(mU/L)\*FPG(mmol/L)] / 22.5 (34).

Peripheral systolic blood pressure (SBP) and diastolic blood pressure (DBP) were measured from the right upper arm (Dinamap CARESCAPE V100, GE Healthcare, Milwaukee, WI) at age 3 (n=639), 4 (n=586), 5 (n=613), and 6 years (n=565) (35). At age 6 years, arterial wall thickness was assessed through carotid intima-media thickness (cIMT) using high resolution B-mode ultrasound while arterial stiffness was assessed through carotid-femoral pulse wave velocity (PWV) by applanation tonometry in 422 and 443 children respectively (Detailed cIMT and PWV protocols in supplementary section).

***Covariates***

Maternal age, ethnicity, educational attainment, household income, parity, and self-reported pre-pregnancy weight were collected at recruitment through interviewer-administered questionnaires. At ~26-28wk of gestation, venous FPG was measured (36). Height was measured using a calibrated stadiometer (SECA213 Stadiometer, SECA Corp, Hamburg, Germany). Pre-pregnancy BMI (ppBMI) was calculated using this height and self-reported pre-pregnancy weight. Plasma cotinine level was measured by liquid chromatography-tandem mass spectrometry (37). Total gestational weight gain (GWG) was calculated from the difference between last measured antenatal weight (taken within 4 weeks from delivery) and self-reported pre-pregnancy weight. Child sex was taken from medical records. Gestational age (in weeks) was calculated based on first trimester crown-rump lengths assessed by trained ultrasonographers. Size at birth was categorized based on birthweight and gestational age using customized birth charts (38). Infants were categorized as low birthweight if they weighed less than 2500g. Paternal height (in cm) was measured during the year 2 postnatal visits.

***Statistical analysis***

To compare differences between participants included and excluded in the study (n=450 families excluded, n=460 children excluded due to 10 pairs of twins), as well as those with and without at least 1 MRI/MRS measurement, blood glucose/insulin measurement, and outcome measurement at year 6, two-tailed t-tests were performed for continuous variables while chi-square tests were performed for categorical variables. Multiple linear regression was performed to test associations between growth groups and childhood cardiometabolic risk biomarkers (anthropometry, BF%, abdominal fat, intramyocellular lipids, liver fat, FPG, fasting insulin, HOMA-IR, cIMT, PWV), with respect to the reference group. To gauge the magnitude of effects, adjusted differences between each growth group and the reference group were also expressed as a percentage change compared to the reference group (except for anthropometry z-scores where negative mean z-scores of the reference group prevent the meaningful evaluation of percentage change). Postestimation pairwise comparisons using Tukey’s adjustment for multiple comparisons were also performed. Based on literature review, the following confounders which influence both fetal/postnatal growth and cardiometabolic outcomes were used: sociodemographic factors (maternal education, child sex and ethnicity), in-utero environmental factors (parity, ppBMI, maternal height, gestational FPG, GWG, cotinine). To determine longitudinal associations of growth groups with SBP and DBP, linear mixed effects (LME) modeling which accounts for correlation between repeated measures in the same participant was performed using maximum likelihood estimation, assuming outcome data was missing at random (39), and defining covariances as unstructured to distinctly estimate all variances and covariances for individual growth curves (40). The models included a random intercept, random linear slope for age, and an age-growth group interaction term, and the same confounders as the regression model above. Using the “margins” command in Stata, adjusted differences in LME-predicted blood pressure between each growth group and the reference group, while holding covariates constant at their mean values, was reported.

A few sensitivity analyses were performed. First, paternal height (available in 78.3% of participants) was added as a covariate in multiple linear regression models, due to its contribution to child growth (41). Since GWG and FPG could potentially be affected by fetal growth due to the temporal overlap in the collection of these variables, we also ran the model after removing GWG and FPG as confounders in multiple linear regression. Standardized residuals of blood pressure regressed on sex and height at each time-point was calculated and their longitudinal associations with the growth groups was evaluated using the same LME model above as blood pressure in growing children is closely related to sex and height. We adjusted for the rate of gestational weight gain from 15-35 weeks of gestation instead of total gestational weight gain, to ensure that this confounder is independent of gestational age. We also additionally adjusted for body fat partitioning at age 4.5 years as it could be a potential mediator of metabolic and cardiovascular outcomes at age 6 years. Due to small sample sizes, we conducted multiple linear regression to determine the main effects of FGD, RPWG, and their interaction term as a sensitivity analysis rather than as a main analysis. Multiple imputation using chained equations was performed to generate 50 imputed datasets for missing covariates (maternal education: n=6, maternal height: n=15, ppBMI: n=63, FPG: n=34, GWG: n=71, cotinine: n=88). The imputation model included all exposure, outcome and confounder variables included in this study (42). Multiple linear regression and LME models, whether including or excluding SGA infants, were performed on imputed datasets where Rubin’s combination rules were used to obtain pooled estimates (43). All analyses were performed using Stata16.0 (StataCorp LP, TX).

# Results

***Cohort description***

(Table 1 here)

(Table 2 here)

Of 1247 participants, 797 were classified into 4 growth groups (Supplementary Fig.S1). Mothers of excluded children had lower education, household income, and age (Supplementary Table S1). Excluded children had lower mean gestational age and birthweight but had similar childhood cardiometabolic outcomes as included offspring (Supplementary Table S2). Among 797 included participants, those with at least 1 MRI or MRS measurement were more likely to have lower maternal education, lower household income, lower proportion of nulliparous mothers, male offspring and Chinese, as well as higher maternal ppBMI and BF% at age 6 years than those without (Supplementary Table S3). Those with at least 1 child blood glucose or insulin measurement had similar parental and child characteristics as those without, except for slightly lower proportion of nulliparous mothers (40.5% vs 49.0%) (Supplementary Table S4). Those with at least 1 outcome measurement at age 6 years had similar parental and child characteristics as those without (Supplementary Table S5).

The mismatch and reference groups had higher maternal education and household income than the FGD-only and RPWG-only groups (Table 1). Compared to all other groups, the mismatch group had the highest proportion of nulliparous mothers (57.6%), mean plasma cotinine levels (4.73±22.12ng/ml), mean height of parents, as well as lowest gestational age (38.67±1.01wk) and birthweight (2.87±0.31kg). Groups which experienced FGD (mismatch, FGD-only) had lower gestational FPG, GWG, and higher proportion of male offspring. Groups which experienced RPWG (mismatch, RPWG-only) had higher proportion of Indians. RPWG-only children had the highest mean z-BMI, z-Height, BF%, and abdominal fat while mismatch children had the highest mean FPG, fasting insulin, HOMA-IR, and blood pressure (Table 2).

***Associations of growth groups with adiposity outcomes***

Compared to the reference group, the mismatch and RPWG-only groups had elevated z-BMI (mismatch: 0.53units [0.19,0.88]; RPWG-only: 0.98units [0.72,1.24]), abdominal subcutaneous adipose tissue (mismatch: 210.30cc [52.74,367.85]; RPWG-only: 315.58cc [196.23,434.94]), and visceral adipose tissue (mismatch: 40.17cc [5.28,75.06]; RPWG-only: 38.03cc [11.60,64.46]). However, only the RPWG-only group had increased overall BF% (4.26% [2.34, 6.19]) while the mismatch group had elevated intramyocellular lipids (0.13% of water signal [0.00,0.26]) and liver fat (0.17% by weight [-0.01,0.35]) (Fig.1A-F, Supplementary Table S6). In fact, the RPWG-only group had noticeably higher z-BMI, BF%, and abdominal subcutaneous adipose tissue than the FGD-only and mismatch group (Supplementary Table S12). The FGD-only group was similar in adiposity outcomes as the reference group (Fig.1A-F, Supplementary Table S6). Relative to the reference group, these findings represent large percentage increases of 45%, 22%, 29%, and 32% in abdominal subcutaneous adipose tissue, visceral adipose tissue, intramyocellular lipids, and liver fat respectively in the mismatch group, compared to a relatively small percentage increase of 4% in BF%. They also represent large percentage increases of 23%, 67%, 21%, and 21% in BF%, abdominal subcutaneous adipose tissue, visceral adipose tissue, and intramyocellular lipids respectively in the RPWG-only group.

***Associations of growth groups with metabolic outcomes***

Compared to the reference group, the mismatch and RPWG-only groups had elevated fasting insulin concentrations (mismatch: 9.04pmol/L [0.61,17.47]; RPWG-only: 8.86pmol/L [3.15,14.57]) and HOMA-IR (mismatch: 0.29units [0.04,0.55]; RPWG-only: 0.28units [0.11,0.45]) without elevated FPG at age 6 years (Fig.1I-K, Supplementary Table S6). The FGD-only group did not have clear alterations in metabolic outcomes compared to the reference group (Fig.1I-K, Supplementary Table S6), and had lower fasting insulin (-10.79pmol/L [-22.14, 0.57]) and HOMA-IR (-0.39units [-0.75, -0.04]) than the mismatch group (Supplementary Table S12). Relative to the reference group, these findings represent large percentage increases of 34%, 33%, 38%, and 36% in fasting insulin (mismatch group), fasting insulin (RPWG-only group), HOMA-IR (mismatch group), and HOMA-IR (RPWG-only group) respectively.

***Associations of growth groups with cardiovascular outcomes***

From age 3-6 years, mismatch children had the highest SBP and DBP LME trajectories (Fig.2). The differences in blood pressure between the mismatch and reference group were more marked at age 3 years and were attenuated at age 6 years: age 3 years (SBP: 6.14mmHg [3.97,8.31]; DBP: 3.04mmHg [1.53,4.54]), 4 years (SBP: 5.13mmHg [3.42,6.83]; DBP: 2.75mmHg [1.58,3.92]), 5 years (SBP: 4.11mmHg [2.41,5.81]; DBP: 2.46mmHg [1.28,3.64]), 6 years (SBP: 3.10mmHg [0.95,5.24]; DBP: 2.18mmHg [0.65,3.70]) (Fig.2, Supplementary Table S7). Similarly, RPWG-only children had elevated SBP (age 3-6 years) and DBP (age 3-5 years) trajectories. The diastolic blood pressure trajectory of the FGD-only children was slightly higher than the reference children consistently from ages 3 to 6 years. The systolic blood pressure trajectory of the FGD-only children started at the same point as the reference children at age 3 years but rose faster than the reference children till it converged with that of the RPWG-only children at age 6 years. None of the groups had substantial alterations in cIMT and PWV, with percentage changes relative to the reference group ranging from -1% to 6% (Fig.1G-H, Supplementary Table S6).

***Sensitivity analysis***

Associations remained similar after additional adjustment for paternal height except that for RPWG-only children, magnitude of association with elevated metabolic outcomes at age 6 years increased slightly (Supplementary Table S8). After removal of GWG and FPG as confounders, associations remained similar except that for the mismatch group, magnitude of association with elevated metabolic outcomes at age 6 years increased slightly (Supplementary Table S9). After using sex- and height-standardized blood pressures, mismatch children still had the highest overall SBP and DBP trajectories from age 3-6 years with the differences attenuated at age 6 years (Supplementary Fig.S2). Associations remained similar after adjusting for rate of gestational weight gain instead of total gestational weight gain, except that association with fasting insulin in the mismatch group increased slightly (Supplementary Table S11). After additional adjustment for body fat partitioning at age 4.5 years, associations of the mismatch and RPWG-only groups with elevated fasting insulin and HOMA-IR was greatly diminished (Supplementary Table S13). Using an interaction model, we found that there is a marked negative interaction between FGD and RPWG for BF% (-4.30% [-7.90, -0.71]), concordant with our finding that a combination of FGD and RPWG was linked to lower BF% than a RPWG-only growth pattern. (Supplementary Table S15).

After performing multiple imputation to account for missing confounders, associations remained similar except that for the mismatch group, magnitude of associations with metabolic outcomes increased: higher FPG (0.13mmol/L [0.01,0.26]; excluding SGA infants: 0.18mmol/L [0.03,0.32]), fasting insulin (11.88pmol/L [4.33,19.44]; excluding SGA infants: 12.54pmol/L [3.41,21.66]), and HOMA-IR (0.44units [0.20,0.68]; excluding SGA infants: 0.48units [0.19,0.77]) (Supplementary Table S10). From LME models after multiple imputation, the mismatch group still had the highest SBP and DBP from 3-6 years old compared to all other growth groups, whether including or excluding SGA infants from the analyses (Supplementary Fig.S3).

# Discussion

Poor in-utero growth and accelerated postnatal growth in the developmentally plastic first 1000 days of life have been linked to adverse cardiometabolic outcomes in adulthood. However, the precise pathways through which early life growth patterns influence later life cardiometabolic outcomes are not fully known as most of the current results have been obtained from retrospective studies in adults. The current study, based on a prospective birth cohort, addresses some of these gaps by investigating how fetal growth deceleration, rapid postnatal weight gains in infancy, or a combination of these two patterns alter cardiometabolic biomarkers in childhood.

We found that rapid weight gain from age 0 to 2 years, whether with or without prior fetal growth deceleration, was associated with a cluster of unfavorable cardiometabolic markers manifesting as early as age 3 years in an Asian cohort. Compared to the reference group which did not experience fetal growth deceleration or rapid postnatal weight gain, the RPWG-only group had elevated BF% (+23%) accompanied by elevated HOMA-IR (36%), subcutaneous adipose tissue (+67%), visceral adipose tissue (+21%) and intramyocellular lipids (+21%). However, it was notable that despite a much lower elevation in BF% (+4%), children in the mismatch group also had elevated HOMA-IR (+38%), subcutaneous adipose tissue (+45%), visceral adipose tissue (+22%) and intramyocellular lipids (+29%). The mismatch group also had a marked elevation in liver fat compared to the RPWG-only group (+32% vs +9%). The associations were not attenuated after excluding SGA children from the analysis, indicating that the above associations are not driven by children with extreme birthweights. The associations of the RPWG-only and mismatch growth patterns with HOMA-IR at age 6y, was lost when we adjusted for the body fat partitioning at age 4.5y, suggesting that fat partitioning may act as a mediator. Our findings also indicate that alterations in fat partitioning and HOMA-IR may emerge even before elevation in adiposity in children with the mismatch growth pattern.

Among all the growth groups, the mismatch group had the highest systolic and diastolic blood pressure trajectories between 3 to 6 years followed by the RPWG group. The systolic blood pressure at age 6 years in the FGD-only children was close to that of the reference group at age 3 years, but was comparable to that of RPWG-only group at age 6 years, despite FGD-only children being leaner and shorter. Long term blood pressure is regulated by the kidney. In humans nephrogenesis (formation of nephrons which are the functional units of the kidney) ceases by 34-36 weeks of gestation. Poor fetal growth in the second and third trimester therefore can influence both nephron endowment and developmental programming of kidney function (44–46). The effects of fetal growth deceleration on blood pressure was amplified with increasing body size (which increases the filtration load) as the children grow older, and with rapid postnatal weight gain (47,48).

Children in both the FGD-only group and the mismatch group experienced downward centile crossing in-utero. This is likely to be a fetal response to prenatal perturbations like maternal undernutrition or placental insufficiency and not due to constitutional factors. However, metabolic characteristics, body fat partitioning and BF% in FGD-only group were very similar to the reference group. Our results suggest that early alterations in the above characteristics in children who experienced fetal growth deceleration require a second hit in the form of rapid postnatal growth. Further follow-up of these children is required to evaluate if the effects of an adverse intrauterine environment are unmasked as they grow older.

Children of all 4 growth groups had similar arterial thickness, and arterial stiffness at age 6y, likely due to small inter-individual variability of these measures in early childhood (49,50). Thus, increase in arterial thickness and stiffness does not seem to play a role in the elevations in blood pressure observed in FGD-only, mismatch and RPWG-only groups. Further longitudinal tracking of these children is required to check if alterations in these cardiometabolic markers emerge later in life and to understand the development of differential cardiometabolic risk.

Although we have detected elevations in cardiometabolic risk markers in the mismatch, FGD-only and RPWG-only group, there is limited evidence to conclude whether these changes are clinically significant in childhood because of the absence of any hard clinical endpoints, which will mainly become evident only in the later stage of life. However, since cardiometabolic risk factors tend to track to adulthood (51,52), children in the RPWG-only, FGD-only and mismatch groups are likely to be at increased risk of adverse cardiometabolic outcomes in adulthood. The alterations in the risk markers in the FGD-only and mismatch group emerged even without increased adiposity. Thus these alterations may be further accentuated if these children become obese later in life. These findings also highlight the importance of monitoring both fetal and childhood growth patterns for early risk stratification, as size at birth or overweight/obesity status alone may not be sufficient to identify all children who may be at elevated cardiometabolic risk.

Our findings were consistent with literature which reported increased subcutaneous adipose tissue, visceral adipose tissue (53,54), total body fat, BF% (13,55), BMI (13,56), and fasting insulin (57) associated with rapid weight gain in childhood. Our findings were also consistent with literature on the adverse cardiometabolic outcomes of poor fetal growth followed by rapid postnatal growth (5,8,9,11–14). For the mismatch group, other studies have reported increased android/gynoid fat ratio, fat mass index, abdominal fat, and BMI at age 6 years (24), increased visceral fat index and liver fat at age 10 years (25), and elevated blood pressure in school-age children and adulthood (12,26). We go on to demonstrate that adverse body fat partitioning alterations manifest as early as age 4.5 years, while elevated blood pressures manifest as early as age 3 years in Asian children.

Our findings contrast with some previous reports which have suggested that accelerated postnatal growth after poor fetal growth may occur with preferential fat accumulation (4,53). The mismatch group, which experienced RPWG after FGD, did not have elevations in overall BF%. This could potentially be due to the way mismatch was characterized. In the current work, we defined mismatch as having slower growth in-utero and accelerated growth in infancy. Older works have used low birth-weight as a proxy for poor fetal growth, which cannot separate the effects of constitutional factors from prenatal perturbations (15). Furthermore, some studies have also noted the absence of excess fat in the mismatch group (58) and reported that catch-up after poor fetal growth might not necessarily involve preferential fat mass accretion (59,60). Our findings suggest that the impact of the mismatch growth pattern might first be on increased ectopic fat accumulation, rather than on increased adiposity. Further investigation is warranted to resolve these conflicting findings on phenomenon of preferential catch-up fat accumulation.

One strength of our study is the use of serial ultrasound measures to measure fetal growth deceleration even if it did not result in extremes of fetal or birth size. While retrospective cohort studies have revealed links between early life growth patterns and cardiometabolic outcomes in adulthood, the precise mechanisms are not fully understood. The extensive phenotyping of cardiometabolic risk markers like body composition, fat partitioning, insulin resistance, blood pressure, arterial thickness and arterial stiffness during childhood in the current study provides insights into the potential pathways through which early life growth patterns could influence later health outcomes. Identifying early cardiometabolic alterations of high-risk growth groups in a large prospective cohort provides evidence for risk-stratification in children and provides impetus to uncover underlying mechanisms.

Our limitations include the fact that our study included only Asian children and excluded preterm infants. Hence, findings from our multi-ethnic Asian cohort might not be generalizable to other populations or to preterm children. Participants with MRI/MRS measures were of slightly lower socioeconomic status and had slightly higher maternal and child adiposity than those without, which might lead to potential bias. We also acknowledge that some children might be classified as having undergone RPWG/FGD due to regression to the mean. Although we have adjusted for sociodemographic and maternal confounders in our analyses, we acknowledge that fetal/postnatal growth and cardiometabolic outcomes might be further influenced by unmeasured behavioral, environmental, nutritional and genetic confounders. While we have suggested possible etiologies underlying our findings, this paper is not focused on the etiologies of the 4 growth groups and further studies are needed to understand them.

# Conclusion

Rapid weight gain from age 0 to 2 years, whether with or without poor fetal growth, is associated with a cluster of elevated cardiometabolic risk markers in early childhood. Notably, children with a mismatch between poor fetal growth and rapid postnatal weight gain had elevated insulin resistance, ectopic fat accumulation and the highest blood pressure trajectory in early childhood, even without a corresponding elevation in BF%. Children with fetal growth deceleration in the absence of rapid postnatal weight gain also had elevated systolic blood pressure without having elevated BF%. The emergence of cardiometabolic alterations even without increased adiposity in the mismatch and FGD-only groups is troubling, as these changes may be further accentuated with later life obesity. Our work highlights the role of longitudinal growth monitoring in both the fetal and postnatal period for early risk stratification, as screening based on size at birth or BMI at a single time point may not be able to uncover all children who may be at increased cardiometabolic risk.

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# Conflict of interest

KMG, PDG and YSC are part of an academic consortium that has received research funding and have received reimbursement for speaking at conferences sponsored by companies selling nutritional products. All other authors declare no potential of interest that might bias the submitted work and no other relationships or activities that could appear to have influenced the submitted work.

# Authors’ contributions

YYO analyzed data, interpreted data, and wrote the manuscript. NM contributed to the analysis, interpretation of data, and drafting of manuscript. SAS, IMA, MTT, WLY, JYH, YHC, SN, SLL, SSV, MVF, KMG, LPS, KHT, PDG, FY, JTC, LHL, KT, LC, NK, YSC, JGE, MEW, SYC, NM, and YSL contributed to the conception and design of the study. IMA, JYH, YHC, and YSL contributed to the analysis and interpretation of data. SAS, IMA, MTT, WLY, JYH, YHC, SN, SLL, SSV, MVF, KMG, LPS, KHT, PDG, FY, JTC, LHL, KT, LC, NK, YSC, JGE, MEW, SYC, and YSL made critical revision of the manuscript for important intellectual content. YYO, NM, and YSL have primary responsibility for the final content.

**References:**

1. Gluckman PD, Hanson MA. Developmental Origins of Disease Paradigm: A Mechanistic and Evolutionary Perspective. Pediatric Research. 2004 Sep;56(3):311–7.

2. Gluckman Peter D., Hanson Mark A., Low Felicia M. Evolutionary and developmental mismatches are consequences of adaptive developmental plasticity in humans and have implications for later disease risk. Philosophical Transactions of the Royal Society B: Biological Sciences. 2019 Apr 15;374(1770):20180109.

3. Godfrey KM, Lillycrop KA, Burdge GC, Gluckman PD, Hanson MA. Epigenetic mechanisms and the mismatch concept of the developmental origins of health and disease. Pediatr Res. 2007 May;61(5 Pt 2):5R-10R.

4. Dulloo AG, Jacquet J, Seydoux J, Montani J-P. The thrifty “catch-up fat” phenotype: its impact on insulin sensitivity during growth trajectories to obesity and metabolic syndrome. Int J Obes (Lond). 2006 Dec;30 Suppl 4:S23-35.

5. Nobili V, Alisi A, Panera N, Agostoni C. Low birth weight and catch-up-growth associated with metabolic syndrome: a ten year systematic review. Pediatr Endocrinol Rev. 2008 Dec;6(2):241–7.

6. Vickers MH, Breier BH, Cutfield WS, Hofman PL, Gluckman PD. Fetal origins of hyperphagia, obesity, and hypertension and postnatal amplification by hypercaloric nutrition. American Journal of Physiology-Endocrinology and Metabolism. 2000 Jul 1;279(1):E83–7.

7. Painter RC, Roseboom TJ, Bleker OP. Prenatal exposure to the Dutch famine and disease in later life: An overview. Reproductive Toxicology. 2005 Sep 1;20(3):345–52.

8. Bavdekar A, Yajnik CS, Fall CH, Bapat S, Pandit AN, Deshpande V, et al. Insulin resistance syndrome in 8-year-old Indian children: small at birth, big at 8 years, or both? Diabetes. 1999 Dec;48(12):2422–9.

9. Berends LM, Dearden L, Tung YCL, Voshol P, Fernandez-Twinn DS, Ozanne SE. Programming of central and peripheral insulin resistance by low birthweight and postnatal catch-up growth in male mice. Diabetologia. 2018;61(10):2225–34.

10. Tarry-Adkins JL, Ozanne SE. Poor Early Growth and Age-Associated Disease. In: Harris JR, Korolchuk VI, editors. Biochemistry and Cell Biology of Ageing: Part II Clinical Science. Singapore: Springer Singapore; 2019. p. 1–19. (Subcellular Biochemistry).

11. Fabricius-Bjerre S, Jensen RB, Færch K, Larsen T, Mølgaard C, Michaelsen KF, et al. Impact of Birth Weight and Early Infant Weight Gain on Insulin Resistance and Associated Cardiovascular Risk Factors in Adolescence. PLoS One. 2011 Jun 2;6(6).

12. Huxley RR, Shiell AW, Law CM. The role of size at birth and postnatal catch-up growth in determining systolic blood pressure: a systematic review of the literature. J Hypertens. 2000 Jul;18(7):815–31.

13. Ong KKL, Ahmed ML, Emmett PM, Preece MA, Dunger DB. Association between postnatal catch-up growth and obesity in childhood: prospective cohort study. BMJ. 2000 Apr 8;320(7240):967–71.

14. Lurbe E, Garcia-Vicent C, Torro MI, Aguilar F, Redon J. Associations of birth weight and postnatal weight gain with cardiometabolic risk parameters at 5 years of age. Hypertension. 2014 Jun;63(6):1326–32.

15. Ananth CV, Vintzileos AM. Distinguishing pathological from constitutional small for gestational age births in population-based studies. Early Hum Dev. 2009 Oct;85(10):653–8.

16. Sonnenschein-van der Voort AMM, Jaddoe VWV, Raat H, Moll HA, Hofman A, de Jongste JC, et al. Fetal and infant growth and asthma symptoms in preschool children: the Generation R Study. Am J Respir Crit Care Med. 2012 Apr 1;185(7):731–7.

17. Bigaard J, Frederiksen K, Tjønneland A, Thomsen BL, Overvad K, Heitmann BL, et al. Body Fat and Fat-Free Mass and All-Cause Mortality. Obesity Research. 2004;12(7):1042–9.

18. Speliotes EK, Massaro JM, Hoffmann U, Vasan RS, Meigs JB, Sahani DV, et al. Fatty liver is associated with dyslipidemia and dysglycemia independent of visceral fat: The Framingham heart study. Hepatology. 2010;51(6):1979–87.

19. Després J-P. Abdominal obesity: the most prevalent cause of the metabolic syndrome and related cardiometabolic risk. Eur Heart J Suppl. 2006 May 1;8(suppl\_B):B4–12.

20. Brumbaugh DE, Crume TL, Nadeau K, Scherzinger A, Dabelea D. Intramyocellular Lipid Is Associated with Visceral Adiposity, Markers of Insulin Resistance, and Cardiovascular Risk in Prepubertal Children: The EPOCH Study. J Clin Endocrinol Metab. 2012 Jul 1;97(7):E1099–105.

21. Soh S-E, Tint MT, Gluckman PD, Godfrey KM, Rifkin-Graboi A, Chan YH, et al. Cohort Profile: Growing Up in Singapore Towards healthy Outcomes (GUSTO) birth cohort study. Int J Epidemiol. 2014 Oct 1;43(5):1401–9.

22. Nesbitt-Hawes EM, Tetstall E, Gee K, Welsh AW. Ultrasound (in)accuracy: it’s in the formulae not in the technique - assessment of accuracy of abdominal circumference measurement in term pregnancies. Australas J Ultrasound Med. 2014 Feb;17(1):38–44.

23. Papageorghiou AT, Ohuma EO, Altman DG, Todros T, Cheikh Ismail L, Lambert A, et al. International standards for fetal growth based on serial ultrasound measurements: the Fetal Growth Longitudinal Study of the INTERGROWTH-21st Project. Lancet. 2014 Sep 6;384(9946):869–79.

24. Gishti O, Gaillard R, Manniesing R, Abrahamse-Berkeveld M, van der Beek EM, Heppe DHM, et al. Fetal and Infant Growth Patterns Associated With Total and Abdominal Fat Distribution in School-Age Children. J Clin Endocrinol Metab. 2014 Jul 1;99(7):2557–66.

25. Vogelezang S, Santos S, Toemen L, Oei EHG, Felix JF, Jaddoe VWV. Associations of Fetal and Infant Weight Change With General, Visceral, and Organ Adiposity at School Age. JAMA Netw Open. 2019 Apr 5;2(4):e192843–e192843.

26. Toemen L, De Jonge LL, Gishti O, Van Osch-Gevers L, Taal HR, Steegers EAP, et al. Longitudinal growth during fetal life and infancy and cardiovascular outcomes at school-age. Journal of Hypertension. 2016;34(7):1396–406.

27. Aris IM, Bernard JY, Chen L-W, Tint MT, Pang WW, Lim WY, et al. Infant body mass index peak and early childhood cardio-metabolic risk markers in a multi-ethnic Asian birth cohort. Int J Epidemiol. 2017 Apr 1;46(2):513–25.

28. WHO Multicentre Growth Reference Study Group. WHO child growth standards: length/height for age, weight-for-age, weight-for-length, weight-for-height and body mass index-for-age, methods and development. Geneva: World Health Organization [Internet]. 2006 [cited 2018 Nov 19]; Available from: http://www.who.int/childgrowth/standards/technical\_report/en/

29. Cole TJ, Green PJ. Smoothing reference centile curves: The lms method and penalized likelihood. Statistics in Medicine. 1992;11(10):1305–19.

30. Chen L-W, Tint M-T, Fortier MV, Aris IM, Shek LP-C, Tan KH, et al. Body composition measurement in young children using quantitative magnetic resonance: a comparison with air displacement plethysmography. Pediatr Obes. 2018;13(6):365–73.

31. Sadananthan SA, Tint MT, Michael N, Aris IM, Loy SL, Lee KJ, et al. Association Between Early Life Weight Gain and Abdominal Fat Partitioning at 4.5 Years is Sex, Ethnicity, and Age Dependent. Obesity. 2019;27(3):470–8.

32. Sadananthan SA, Prakash B, Leow MK-S, Khoo CM, Chou H, Venkataraman K, et al. Automated segmentation of visceral and subcutaneous (deep and superficial) adipose tissues in normal and overweight men. Journal of Magnetic Resonance Imaging. 2015 Apr 1;41(4):924–34.

33. Michael N, Gupta V, Sadananthan SA, Sampathkumar A, Chen L, Pan H, et al. Determinants of Intramyocellular liquid accumulation in early childhood. International Journal of Obesity. 2019 Jul 3;

34. Matthews DR, Hosker JP, Rudenski AS, Naylor BA, Treacher DF, Turner RC. Homeostasis model assessment: insulin resistance and β-cell function from fasting plasma glucose and insulin concentrations in man. Diabetologia. 1985 Jul 1;28(7):412–9.

35. Aris IM, Bernard JY, Chen L-W, Tint MT, Lim WY, Soh SE, et al. Postnatal height and adiposity gain, childhood blood pressure and prehypertension risk in an Asian birth cohort. International Journal of Obesity. 2017 Jul;41(7):1011–7.

36. Aris IM, Soh SE, Tint MT, Saw SM, Rajadurai VS, Godfrey KM, et al. Associations of gestational glycemia and prepregnancy adiposity with offspring growth and adiposity in an Asian population. Am J Clin Nutr. 2015 Nov 1;102(5):1104–12.

37. Ng S, Aris IM, Tint MT, Gluckman PD, Godfrey KM, Shek LP-C, et al. High Maternal Circulating Cotinine During Pregnancy is Associated With Persistently Shorter Stature From Birth to Five Years in an Asian Cohort. Nicotine Tob Res. 2019 Jul 17;21(8):1103–12.

38. Mikolajczyk RT, Zhang J, Betran AP, Souza JP, Mori R, Gülmezoglu AM, et al. A global reference for fetal-weight and birthweight percentiles. The Lancet. 2011 May 28;377(9780):1855–61.

39. Finucane MM, Samet JH, Horton NJ. Translational methods in biostatistics: linear mixed effect regression models of alcohol consumption and HIV disease progression over time. Epidemiol Perspect Innov. 2007 Sep 19;4:8.

40. Johnson W, Balakrishna N, Griffiths PL. Modeling physical growth using mixed effects models. Am J Phys Anthropol. 2013 Jan;150(1):58–67.

41. Griffiths LJ, Dezateux C, Cole TJ. Differential parental weight and height contributions to offspring birthweight and weight gain in infancy. Int J Epidemiol. 2007 Feb 1;36(1):104–7.

42. White IR, Royston P, Wood AM. Multiple imputation using chained equations: Issues and guidance for practice. Statistics in Medicine. 2011;30(4):377–99.

43. Rubin DB. Multiple imputation for nonresponse in surveys. Hoboken, N.J: Wiley-Interscience; 2004. 287 p. (Wiley classics library).

44. Verburg BO, Geelhoed JJM, Steegers EAP, Hofman A, Moll HA, Witteman JCM, et al. Fetal kidney volume and its association with growth and blood flow in fetal life: The Generation R Study. Kidney International. 2007 Sep 2;72(6):754–61.

45. Bakker H, Gaillard R, Franco OH, Hofman A, van der Heijden AJ, Steegers EAP, et al. Fetal and infant growth patterns and kidney function at school age. J Am Soc Nephrol. 2014 Nov;25(11):2607–15.

46. Luyckx VA, Brenner BM. The Clinical Importance of Nephron Mass. JASN. 2010 Jun 1;21(6):898–910.

47. Correia-Costa L, Schaefer F, Afonso AC, Bustorff M, Guimarães JT, Guerra A, et al. Normalization of glomerular filtration rate in obese children. Pediatr Nephrol. 2016 Aug 1;31(8):1321–8.

48. Brandt JR, Wong CS, Hanrahan JD, Qualls C, McAfee N, Watkins SL. Estimating absolute glomerular filtration rate in children. Pediatr Nephrol. 2006 Dec;21(12):1865–72.

49. Hvidt KN. Blood pressure and arterial stiffness in obese children and adolescents. Dan Med J. 2015 Mar;62(3).

50. Park MH, Skow Á, De Matteis S, Kessel AS, Saxena S, Viner RM, et al. Adiposity and carotid-intima media thickness in children and adolescents: a systematic review. BMC Pediatr. 2015 Oct 16;15:161.

51. Juhola J, Magnussen CG, Viikari JSA, Kähönen M, Hutri-Kähönen N, Jula A, et al. Tracking of Serum Lipid Levels, Blood Pressure, and Body Mass Index from Childhood to Adulthood: The Cardiovascular Risk in Young Finns Study. The Journal of Pediatrics. 2011 Oct 1;159(4):584–90.

52. Camhi SM, Katzmarzyk PT. Tracking of cardiometabolic risk factor clustering from childhood to adulthood. International Journal of Pediatric Obesity. 2010 Jan 1;5(2):122–9.

53. Ibáñez L, Ong K, Dunger DB, de Zegher F. Early Development of Adiposity and Insulin Resistance after Catch-Up Weight Gain in Small-for-Gestational-Age Children. J Clin Endocrinol Metab. 2006 Jun 1;91(6):2153–8.

54. Ibáñez L, Suárez L, Lopez-Bermejo A, Díaz M, Valls C, de Zegher F. Early Development of Visceral Fat Excess after Spontaneous Catch-Up Growth in Children with Low Birth Weight. J Clin Endocrinol Metab. 2008 Mar 1;93(3):925–8.

55. Chomtho S, Wells JC, Williams JE, Davies PS, Lucas A, Fewtrell MS. Infant growth and later body composition: evidence from the 4-component model. Am J Clin Nutr. 2008 Jun 1;87(6):1776–84.

56. Zheng M, Lamb KE, Grimes C, Laws R, Bolton K, Ong KK, et al. Rapid weight gain during infancy and subsequent adiposity: a systematic review and meta-analysis of evidence. Obes Rev. 2018 Mar;19(3):321–32.

57. Voerman E, Jaddoe VWV, Franco OH, Steegers EAP, Gaillard R. Critical periods and growth patterns from fetal life onwards associated with childhood insulin levels. Diabetologia. 2017;60(1):81–8.

58. Milovanovic I, Njuieyon F, Deghmoun S, Chevenne D, Levy-Marchal C, Beltrand J. SGA Children with Moderate Catch-Up Growth Are Showing the Impaired Insulin Secretion at the Age of 4. PLoS One. 2014 Jun 30;9(6).

59. Raaijmakers A, Jacobs L, Rayyan M, van Tienoven TP, Ortibus E, Levtchenko E, et al. Catch-up growth in the first two years of life in Extremely Low Birth Weight (ELBW) infants is associated with lower body fat in young adolescence. PLoS ONE. 2017;12(3):e0173349.

60. van de Lagemaat M, Rotteveel J, Lafeber HN, van Weissenbruch MM. Lean mass and fat mass accretion between term age and 6 months post-term in growth-restricted preterm infants. Eur J Clin Nutr. 2014 Nov;68(11):1261–3.