

REVIEW

Relationship between prenatal growth, postnatal growth and childhood obesity: a review

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Evidence is accumulating that obesity risks become established early in life. The goal of this systematic review is to assess whether there is evidence that rapid postnatal growth holds different implications for childhood obesity risks for children who experienced different patterns of prenatal growth. We conducted systematic database searches in PubMed and Embase in October 2014 for studies assessing the implications of prenatal and postnatal growth for childhood obesity. The 18 studies meeting inclusion criteria indicated that risks of obesity increased with birthweight; risks of obesity also increased with rapid postnatal growth for children across the birthweight distribution. Fifteen studies indicated that rapid postnatal growth is linearly associated with obesity for children and adolescents of all sizes at birth, with no interaction effect. Three studies reported interaction effects with postnatal growth, conferring additional increased risk of obesity among children and adolescents who were small at birth. Both prenatal and postnatal growth are important risk factors for obesity, and their combined effects should be analyzed further to understand how obesity risks develop early in life.

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INTRODUCTION

Obesity is a public health concern worldwide, with the global prevalence of childhood overweight and obesity estimated at 6.7% in 2010.¹ There is mounting evidence that the path to obesity is established early in life, perhaps as early as prenatally and in infancy.^{2–5} What the early life factors are and how they may work together to shape a child's long-term weight trajectory are important areas of ongoing research.

Studies have indicated that *prenatal growth* patterns may influence body fat distribution and the development of obesity later in life.^{4,6–8} It has been hypothesized that the relation between birthweight and obesity may follow a U or J-shaped curve, with higher risk at both extremes of the birthweight distribution.^{7,9} Empirical evidence has shown that obesity risks typically increase linearly with weight at birth: high birthweight infants are at higher risk of long-term overweight;⁸ evidence of higher risk of obesity at the low end of the birthweight distribution has been less frequently observed.^{8,10–12} High birthweight may be associated with later obesity due to larger size stemming from high levels of lean mass,^{13,14} while low birthweight may be associated with higher central adiposity.^{15–18} Studies have also indicated that *postnatal growth* during the first 1–2 years of life may be positively associated with subsequent obesity in childhood, adolescence and adulthood, as obesity risks, both in terms of body mass index (BMI) and adverse body composition, are higher for those who experienced rapid postnatal weight gain.^{19–23}

An important consideration in understanding the early life emergence of obesity risks is the relationship between prenatal and postnatal growth and whether some combinations of growth increase risks. Prenatal growth has been shown to be inversely associated with postnatal growth, with babies born smaller experiencing higher growth during infancy compared with larger babies.^{21,24,25} It has been suggested that babies who experience

growth restriction *in utero* and therefore are born small compensate by growing rapidly in early infancy, and that this *mismatch* between prenatal and postnatal growth may elevate risks of obesity and other chronic diseases.^{26–28} If this is the case, there could be trade-offs between the short-term benefits of 'catch-up' growth and chronic disease risk in the longer-term.

Because most studies that have investigated the relationship between prenatal growth or postnatal growth and obesity have not taken both into account, the relative importance of prenatal and early postnatal growth for obesity risk is still unclear, and the idea of mismatch between prenatal and postnatal growth is difficult to assess. It is also unclear whether rapid postnatal growth confers additional risk across the entire birthweight spectrum, or whether rapid postnatal growth has different implications for obesity risk depending on size at birth. The clinical implications of understanding the relationship between postnatal growth and obesity across the birthweight spectrum are critical to efforts to ensure healthy growth in childhood.

The intention of this systematic review is to elucidate which patterns of prenatal and postnatal growth may be linked to childhood obesity, synthesizing evidence from studies that measured the interaction between prenatal and postnatal growth and their associations with obesity in childhood or adolescence. We summarize and assess evidence for the hypotheses that prenatal growth and obesity are associated independent of postnatal growth, that postnatal growth and obesity are associated independent of prenatal growth, and that interactions exist between prenatal and postnatal growth and subsequent obesity.

DATA AND METHODS

Literature searches were conducted in PubMed and Embase databases in October 2014 using combinations of the following

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search terms for whole articles: *birthweight/low birthweight/small for gestational age AND rapid growth/accelerated growth/catch up growth/catch up weight/rapid postnatal growth/rapid weight gain/rapid infant weight gain/rapid neonatal weight gain/growth rate/growth velocity/infant growth/infant weight gain AND obesity/overweight/adiposity*. Additional studies were sought from citations listed in the retrieved papers and relevant review articles.

Abstracts were reviewed for presentation of original research (no review articles or commentaries) that assessed the interaction between prenatal and postnatal growth measures on subsequent obesity or adiposity in humans prior to age 21. Studies that assessed the independent associations between prenatal or postnatal growth and subsequent obesity but did not consider the interaction between them were excluded. Studies were excluded if full articles were not published in English.

We reviewed full articles if the abstracts met the inclusion criteria or if a decision could not be made from the abstract alone. Information regarding study design, population, sample size, methods, results and limitations was recorded from each article using a standardized abstraction form. The literature screening process is displayed in Figure 1. The study was reported following PRISMA guidelines.

Two authors independently assessed the quality of all studies that met inclusion criteria using the Newcastle-Ottawa Scale.²⁹ The scale addresses three categories: Selection, Comparability and Outcome. Each category includes subitems that are assessed and scored for each study. The Selection category assesses four items, with a maximum score of four points: representativeness of the exposed cohort, selection of the non-exposed cohort, ascertainment of exposure and demonstration that the outcome of interest was not present at the start of the study. The Comparability category has one item: whether the study controls for confounding variables, with a maximum score of two points. The Outcome category assesses outcome ascertainment based on three items, with a maximum score of three points: assessment of the outcome, length of follow-up and adequacy of follow-up. Thus, a study is given one point for each item meeting criteria in the Selection and Outcome categories and two points for meeting criteria in the Comparability category, with a possible range between 0 and 9. Studies achieving 67% or higher (greater than 6 points) are considered 'good' quality; 34–66% (3–6 points) 'fair'; and below 33% (below 3 points) 'poor'.³⁰

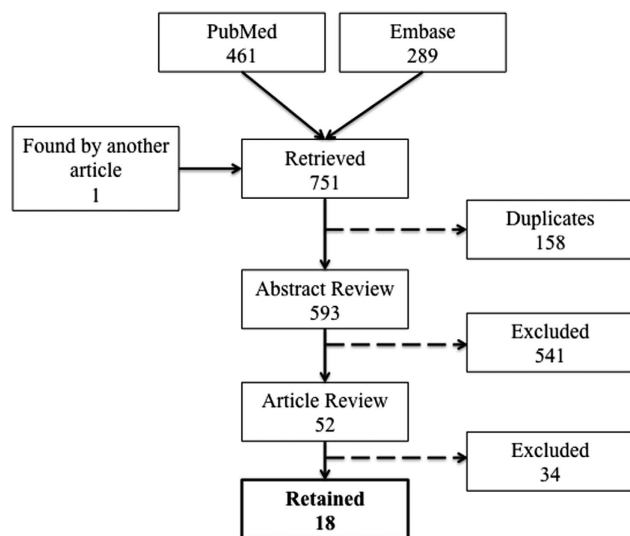


Figure 1. Literature screening for studies assessing the interaction between prenatal and postnatal growth. Numbers indicate article counts retained or excluded at each step of the process.

RESULTS

Characteristics of the 18 publications meeting the inclusion criteria, describing 16 studies, are shown in Table 1. All 18 publications reported results from cohort studies, 16 of which were prospective and 8 of which were population based. Study participants were born between 1959 and 2006. Four reports were published in 2000–2005, eight in 2006–2010 and six in 2011–2014. Nine (50%) reports were based on data from Europe, five (28%) from North America and four (22%) from other regions. Sample size ranged from 163 to 19 397: seven reports (39%) had 150–500 participants, three (17%) had 500–1000 participants, six (33%) had 1000–10 000 participants and two (11%) had >10 000 participants. One study excluded infants weighing <2500 g at birth; nine studies excluded infants born preterm (gestational age <37 weeks).

The quality of the 18 studies included in this systematic review according to the Newcastle-Ottawa Scale is presented in Table 2.^{31–48} The mean total score was 8.7 out of a maximum score of 9 (range 7–9), indicating that, overall, the methodological quality was high. All 18 studies were awarded the highest possible score (2 stars) for the Comparability category, and 16 out of 18 studies were awarded the highest possible score (3 stars) for the Outcome category, with 2 studies receiving no stars for adequacy of follow-up. For the Selection category, two studies were awarded no stars for representativeness of the exposed cohort, and one study was awarded no stars for ascertainment of exposure.

Measures of prenatal growth

Birthweight is the most commonly used proxy measure of prenatal growth and is used as an indicator of fetal growth restriction when other indicators are not available. Low birthweight (LBW) is generally defined as birthweight <2500 g, high birthweight as >4000 g and normal birthweight as 2500–4000 g.

Of the 18 studies included in this review, 12 used birthweight as a proxy of prenatal growth, with 4 using birthweight in grams or kilograms as a continuous variable, 3 using birthweight categories (quartiles or tertiles) and 5 using z-scores for birthweight⁴ or birthweight for length.¹ Two additional studies used age- and sex-specific z-scores for BMI at birth relative to a reference population. Four other publications considered birthweight in the context of gestational age: three used size-for-gestational-age categories, with small for gestational age (SGA) defined as birthweight ≤5th or ≤10th percentile and large for gestational age (LGA) defined as birthweight ≥90th or ≥95th percentile compared with a referent appropriate for gestational age (AGA) group; the fourth measured intrauterine growth restriction (IUGR), defined as birthweight for gestational age <10th percentile of the Williams curve, a US population-based reference for fetal growth which includes measurements from 20 weeks gestation onward.^{44,49}

One publication further classified LBW infants based on body proportionality using the Ponderal Index (PI) (weight (g)/length (cm)³ × 100) to distinguish between newborns who were stunted (length at birth <−2 z-score and PI ≥2.5) vs wasted (length at birth ≥−2 z-score and PI <2.5).³⁸

Measures of postnatal growth

Standard definitions for rates of growth in the first two years after birth have not been adopted and studies used varying terms and measurements, most commonly 'catch-up growth', 'rapid growth' and 'accelerated growth'. Postnatal growth has commonly been measured as change in weight z-scores or standard deviation (s.d.), often using >0.67 s.d. units. This cutoff was proposed by Ong *et al.*²⁴ to distinguish 'catch-up' growth from normal growth, defined as changes within the range of −0.67 to 0.67 s.d. This cutoff is used because it represents the width of each percentile

Table 1. Design characteristics of studies examining the interaction of prenatal and postnatal growth (*n* = 18)

Reference and (data set name)	Location	n	Study design	Study population	Prenatal growth measure/proxy	Postnatal growth measure	Timing of growth	Obesity outcomes	Age at assessment	Interaction analysis
Chomtho <i>et al.</i> ³¹	London, England	234	Retrospective cohort	Born at term approx. 1988–2004 ^a	BW (s.d. score)	Δ weight s.d. score	Birth–3 mos	BMI FFMI WC	4–20 yrs	Interaction term
Durmus <i>et al.</i> ³² (Generation R Study)	Rotterdam, The Netherlands	481	Population-based prospective cohort	Born 2002–2006	BW (tertiles)	Catch-up growth (increase weight s.d. score > 0.67)	Birth–2 yrs	Ratio of preperitoneal to subcutaneous distance ^b	2 yrs	Stratified analysis
Ekelund <i>et al.</i> ³³ (SWEDES Study)	Stockholm, Sweden	248	Prospective cohort	Born 1984–1986	BW (kg)	Δ weight s.d. score Rapid weight gain (increase weight s.d. score > 0.67)	Birth–6 mos	BMI WC Relative fat mass (%) Fat mass (kg) Fat-free mass (kg)	17 yrs	Interaction term
Eriksson <i>et al.</i> ³⁴ (COMPASS Study)	Stockholm, Sweden	2453	Population-based retrospective cohort	Born 1985–1987	BW (z-score)	Weight z-score	Birth–1 yr	BMI FFMI Fat mass (%) WC	15 yrs	Interaction term
Euser <i>et al.</i> ³⁵ (POPS Study)	The Netherlands	403	Prospective cohort	Born 1983 < 32 wks gestation and/or BW < 1500 g	BW (s.d. score)	Δ weight s.d. score	Birth–3 mos	BMI WC Sum of 4 SFT ^c Fat mass (kg) Fat-free mass (kg) % body fat	19 yrs	Interaction term
Gishti <i>et al.</i> ³⁶ (Generation R Study)	Rotterdam, The Netherlands	6464	Population-based prospective cohort	Born 2002–2006	Size for gestational age: SGA (< 10th pc), AGA (≥ 10th–90th pc), LGA (> 90th pc)	Growth acceleration (ΔSDS score > 0.67) Growth deceleration (ΔSDS score < 0.67)	Birth–2 yrs	BMI FFMI FFMI Android/gynoid fat ratio ^d	6 yrs	Interaction term & stratified analysis
Gomez-Lopez <i>et al.</i> ³⁷ (QUALITY Study)	Quebec, Canada	423	Prospective cohort	Term AGA born approx. 1995–2000 with parental history of obesity	BMI z-score at birth	Δ BMI z-score	Birth–2 yrs	BMI z-score WC Waist-to-height ratio FFMI FFMI % body fat % central to total fat	8–10 yrs	Interaction term
Goncalves <i>et al.</i> ³⁸	Pernambuco, Brazil	167	Prospective cohort	Term-born 1993–1994	BW (z-score)	Catch-up growth (increase weight s.d. score > 0.67)	Birth–6 mos	BMI WC	8 yrs	Stratified analysis
Holzhauser <i>et al.</i> ³⁹ (Generation R Focus Study)	Rotterdam, The Netherlands	818	Population-based prospective cohort	Term-born 2003–2005	BW (s.d. score)	Δ weight s.d. score	Birth–6 mos.	Sum of 4 SFT % body fat Central fat ratio ^e	6 mos	Interaction term
Hui <i>et al.</i> ⁴⁰	Hong Kong, China	6075	Population-based prospective birth cohort	Term-born 1997	BW (tertiles)	Δ weight z-score tertiles Accelerated growth (Δ weight z-score > 0.67)	Birth–3 mos	BMI z-score OW ^f	7 yrs	Interaction term & stratified analysis
Jones-Smith <i>et al.</i> ⁴¹	Morelos, Mexico	163	Prospective cohort	Term-born 1998–2001	BMI z-score at birth	Δ BMI z-score Accelerated growth (positive change in BMI z-score)	Birth–1 yr	BMI z-score OW ^g	4–6 yrs	Interaction term
Jones-Smith <i>et al.</i> ⁴²	Morelos, Mexico	586	Prospective cohort	Term-born 1998–2001	BW	BMI velocity relative to the study population's average growth	Birth–2 yrs	OW ^h	8 yrs	Interaction term

Table 1. (Continued)

Reference and (data set name)	Location	n	Study design	Study population	Prenatal growth measure/proxy	Postnatal growth measure	Timing of growth	Obesity outcomes	Age at assessment	Interaction analysis
Kramer <i>et al.</i> ⁴³ (PROBIT)	Republic of Belarus	13 879	Prospective cohort	Term-born ≥2500 g in 1996–1997	Size for gestational age: SGA (<10th pc), AGA (≥10th–90th pc), LGA (>90th pc)	Catch-up growth (increase weight-for-age z-score >0.67 among SGA children) Catch-down growth (decrease weight-for-age z-score >0.67 among LGA children)	Birth–3 or 6 mos	BMI % body fat FFMI WC Waist-to-hip ratio Subscapular to triceps SFT ratio ^l	11.5 yrs	Stratified analysis
Monteiro <i>et al.</i> ⁴⁴	Pelotas, Brazil	1076	Population-based prospective cohort	Born in 1982 in urban areas	IUGR (BW for gestational age <10th pc)	Rapid growth (Δ weight-for-age z-score >0.67)	Birth–2 yrs.	OW ^j Obesity ^k	14–16 yrs	Interaction term & stratified analysis
Stettler <i>et al.</i> ⁴⁵	Seychelles	5514	Population-based prospective cohort	Born in 1984–1994	BW (quartiles)	Δ weight (kg)	Birth–1 yr	OW ^f Obesity ^f	4.5–17.4 yrs	Stratified analysis
Stettler <i>et al.</i> ⁴⁶	12 US cities	19 397	Prospective cohort study	Term-born 1959–1965	BW (g)	Rate of weight gain (100 g/mo)	Birth–4 mos	OW ^g	7 yrs	Stratified analysis
Taal <i>et al.</i> ⁴⁷	The Netherlands	3941	Population-based prospective cohort	Born 2002–2006	Size for gestational age: SGA (<5th pc) AGA (5th–95th pc) LGA (>95th pc)	Catch-up growth (increase Δ weight s.d. >0.67) Catch-down growth (decrease Δ weight s.d. >0.67)	Birth–2 yrs	BMI s.d. score OW or obesity ^f Sum of 4 SFT Central fat mass ^l Peripheral fat mass ^m	2–4 yrs	Stratified analysis
Taveras <i>et al.</i> ⁴⁸ (Project Viva Study)	Massachusetts, USA	559	Prospective cohort	Born 1999–2003	BW for length z-score	Δ weight-for-length z-score quartiles	Birth–6 mos	BMI z-score Sum of subscapular & triceps SFT Obesity ⁿ	3 yrs	Interaction term & stratified analysis

Abbreviations: AGA, appropriate for gestational age; BMI, body mass index; BW, birthweight; FFMI, fat-free mass index (kg/m²); FMI, fat mass index (kg/m²); IUGR, intrauterine growth restricted; kg, kilograms; LGA, large for gestational age; n, sample size; OW, overweight; pc, percentile; s.d., standard deviation; SFT, skinfold thickness; SGA, small for gestational age; WC, waist circumference; wks, weeks; yrs, years; WHR, waist-to-hip ratio; Δ, change; mos, months. ^aApproximate birth year range calculated backward from publication date. ^bRatio of prepertoneal to subcutaneous distance refers to the ratio between prepertoneal fat thickness and subcutaneous thickness, with a higher ratio representing a more adverse distribution of abdominal fat. ^cSum of four SFT refers to the sum of biceps, triceps, suprailliacal and subscapular fat thicknesses. ^dAndroid/gynoid fat ratio reflects central body fat distribution in the abdomen and hip regions, respectively, and is used as a measure of waist-to-hip fat distribution. ^eCentral fat ratio refers to a ratio of the sum of suprailliacal and subscapular SFT to the sum of triceps and biceps SFT. ^fInternational Obesity Task Force (IOTF) reference for obesity cutoffs. ^gCenters for Disease Control and Prevention (CDC) reference for obesity cutoffs. ^hWorld Health Organization (WHO) reference for obesity cutoffs. ⁱSubscapular to triceps SFT ratio is used to assess regional differences in subcutaneous fat distribution, with a higher ratio representing increased central fat deposition. ^jUS National Health and Nutrition Examination Survey (NHANES) reference for obesity cutoffs. ^kOverweight and tricipital and subscapular SFT in the 90th pc. ^lCentral fat mass refers to the sum of suprailliacal and subscapular SFT. ^mPeripheral fat mass refers to the sum of triceps and biceps SFT.

Table 2. Quality assessment of the 18 studies included in the systematic review according to the Newcastle-Ottawa Scale

	Selection			Outcome not present at baseline	Comparability Analysis adjusted for confounding factors	Outcome			Total score
	Exposed cohort representativeness	Selection of non-exposed cohort	Ascertainment of exposure			Assessment of outcome	Length of follow-up	Adequacy of follow-up	
Chomtho <i>et al.</i> ³¹	0	1	0	1	2	1	1	1	7
Durmus <i>et al.</i> ³²	1	1	1	1	2	1	1	1	9
Ekelund <i>et al.</i> ³³	1	1	1	1	2	1	1	1	9
Eriksson <i>et al.</i> ³⁴	1	1	1	1	2	1	1	1	9
Euser <i>et al.</i> ³⁵	1	1	1	1	2	1	1	1	9
Gishti <i>et al.</i> ³⁶	1	1	1	1	2	1	1	1	9
Gomez-Lopez <i>et al.</i> ³⁷	1	1	1	1	2	1	1	1	9
Goncalves <i>et al.</i> ³⁸	0	1	1	1	2	1	1	1	8
Holzhauser <i>et al.</i> ³⁹	1	1	1	1	2	1	1	1	9
Hui <i>et al.</i> ⁴⁰	1	1	1	1	2	1	1	1	9
Jones-Smith <i>et al.</i> ⁴¹	1	1	1	1	2	1	1	1	9
Jones-Smith <i>et al.</i> ⁴²	1	1	1	1	2	1	1	1	9
Kramer <i>et al.</i> ⁴³	1	1	1	1	2	1	1	0	8
Monteiro <i>et al.</i> ⁴⁴	1	1	1	1	2	1	1	1	9
Stettler <i>et al.</i> ⁴⁵	1	1	1	1	2	1	1	1	9
Stettler <i>et al.</i> ⁴⁶	1	1	1	1	2	1	1	1	9
Taal <i>et al.</i> ⁴⁷	1	1	1	1	2	1	1	0	8
Taveras <i>et al.</i> ⁴⁸	1	1	1	1	2	1	1	1	9

band or the distance between two percentile bands on a growth chart, and it has also been used in recent years to define 'accelerated' or 'rapid' growth relative to normal growth.

Studies in this review used eight measures of postnatal growth, six using change in weight s.d. scores and eight using 0.67 as a cutoff for change in weight z-scores (six studies) and/or change in weight-for-age z-scores (three studies). Other measures were rate of weight gain per month in 100-g increments (one study), change in weight in kilograms (one study), change in BMI z-score (two studies) and change in weight-for-length z-score quartiles (one study). Finally, one study measured postnatal growth using BMI velocity relative to the study population's average growth curve, and another used weight z-score at age 1 year adjusted for birthweight z-score.

The term 'catch-up growth' has been used more consistently than other terms to reflect a change in weight z-score or weight-for-age z-score greater than 0.67 s.d. units; still, studies have differed in who can, by definition, experience catch-up growth. Originally, the term was applied exclusively to children who were low birthweight or small for gestational age, with the idea that these children were catching up to their intended growth curve after restricted intrauterine growth. Although children born small are more likely to experience growth that crosses centile bands upwards on a growth chart, children born at normal birthweight or appropriate for gestational age and children born at the high end of the birthweight distribution or large for gestational age can also cross centile bands. Some studies have expanded the use of the term catch-up growth to describe growth of any child who crosses centile bands upward in the first one or two years of life. Some studies also measured 'catch-down growth', representing a decrease in z-score greater than -0.67 s.d. units. The term catch-down growth has mostly been used to describe slow growth of children who were large for gestational age.

Studies included in this review assessed postnatal growth over varying time periods, most commonly between birth and 1 year (three studies) or birth and 2 years (six studies). Some used shorter time periods: birth to 3 months (three studies), to 4 months (one study) and to 6 months (four studies), and some took measures at multiple time points before age 1 year (nine studies).

Measures of obesity

The 18 publications meeting inclusion criteria used 19 obesity-related outcomes, with one to seven outcomes per study. The most commonly measured outcomes were BMI (12 studies), waist circumference (7 studies), and overweight or obesity based on cutoff points relative to various growth references (8 studies). BMI and cutoffs based on it are a commonly used measure of excess weight but do not account for body fat distribution, fat mass or lean mass. Their validity for use in children has been debated because their accuracy varies with degree of body fatness, having high accuracy as an indicator of excess body fat among obese children and poorer accuracy among overweight and thin children.⁵⁰ Some studies also examined direct measures of adiposity: percent body fat (four studies), fat mass (three studies), fat-free mass (two studies), sum of skinfolds (four studies), relative fat mass (one study), subscapular to triceps skinfold ratio (one study), central fat ratio (one study), central to total fat ratio (one study), waist-to-hip ratio (one study), waist-to-height ratio (one study), ratio of preperitoneal distance to subcutaneous transverse distance (one study), android/gynoid fat ratio (one study), fat mass index (FMI) (five studies) and fat-free mass index (FFMI) (four studies). These measures were collected between the ages of 6 months³⁹ and 20 years,³² with three studies at ages 1–5 years; seven studies at 6–12 years; four studies at 13–19 years; and three studies across age groups.^{31,41,45}

Summary of findings

Independent associations between prenatal growth, postnatal growth and obesity. Ten studies eligible for this review assessed the relation between prenatal and postnatal growth (see Appendix Table A1), and nine of these found that children and adolescents born lighter were more likely to experience rapid growth in early life than children and adolescents born heavier; one study examining birthweight adjusted for length found that heavier newborns tended to have larger changes in weight for length between birth and 6 months.⁴⁸ Since prenatal and postnatal growth patterns are correlated, it is important to understand the independent associations that each may have with obesity.

Studies not adjusting for postnatal growth have reported that birthweight is positively associated with BMI and with odds of obesity in childhood and adolescence.^{4,7,51} In this analysis we report on the relationship between prenatal growth and subsequent obesity after accounting for postnatal growth (estimates shown in six studies) (see Table 3). The majority reported positive associations between prenatal growth and obesity measures even after accounting for postnatal growth. Four studies reported that birthweight or BMI at birth was significantly and positively associated with at least one obesity outcome, with larger newborns having higher measures of BMI (4/4), waist circumference (3/3), relative fat mass (1/1), fat-free mass or fat-free mass index (2/2), and waist-to-height ratio (1/1). On the other hand, two of the six studies reported significant negative associations between birthweight and some measures of obesity after accounting for postnatal growth, with lighter newborns having higher measures of sum of skinfolds (1/1), percent body fat or percent fat mass (2/3) and central fat ratio (1/2). The six studies showed no clear pattern between proxies of prenatal growth and odds of overweight or obesity, central fat ratio, or fat mass or fat mass index. The study that examined boys and girls separately reported that birthweight was positively associated with BMI, waist circumference, and fat-free mass for boys only, and that birthweight was negatively associated with percent body fat for girls only, with lighter newborn girls having a higher percentage of body fat compared with heavier newborn girls.³⁴

Studies not adjusting for prenatal growth have reported that postnatal growth is positively associated with subsequent obesity.^{19,20} In this analysis we report on the relationship between postnatal growth and subsequent obesity after accounting for prenatal growth (estimates shown in 11 studies) (see Table 3). All 11 studies found postnatal growth to be significantly positively associated with at least one measure of obesity: BMI (9/9), overweight or obesity (5/5), waist circumference (6/6), fat mass or fat mass index (6/6), fat-free mass or fat-free mass index (6/6), percent body fat or percent fat mass (4/4), relative fat mass (1/1), central fat ratio (2/2), waist-to-height ratio (1/1) and sum of skinfolds (2/3). These associations were observed in diverse populations across a variety of periods of growth and a range of ages at outcome assessment. They collectively indicate that postnatal growth is significantly associated with obesity independently of prenatal growth.

Interactions between prenatal growth and postnatal growth and obesity. Eighteen studies have reported on the interactions between prenatal and postnatal growth, assessing whether the implications of postnatal growth for obesity are different for children and adolescents who experienced different prenatal growth patterns. These studies used regression models with interaction terms (12 studies) and stratified analyses (10 studies) (see Table 3). Of the 18 studies, 15 reported that postnatal growth confers higher risk of obesity for children and adolescents of all sizes at birth. Four of the 15 studies reported that the interaction terms were not significant but did not show any further results, five reported only the *P*-values for the interaction terms, and six provided coefficient estimates.

Three studies found significant interactions between prenatal and postnatal growth variables, indicating that the relationship between postnatal growth and obesity differs for children according to their prenatal growth. Hui *et al.*,⁴⁰ one of the largest studies, with data from China, examined birthweight tertiles, change-in-weight z-score between birth and 3 months, and BMI and overweight at age 7 years. The interaction between birthweight and change-in-weight z-score was statistically significant ($P=0.001$) and indicated that accelerated growth was more strongly associated with higher BMI in children with lower birthweight than children with higher birthweight. Children in the

lowest birthweight z-score tertile (mean birthweight of 2.8 kg) who experienced accelerated growth (change-in-weight z-score >0.67 s.d.) had similar increases in BMI z-scores by age 7 to children with medium birthweight (mean 3.2 kg) and medium growth (second tertile) and to children with high birthweight (mean 3.6 kg) and slow growth (first tertile) (Δ BMI z-score 0.71, 0.70 and 0.70). The interaction was more pronounced among boys, while girls' weight tracked more with their birthweight than did boys', whatever their postnatal growth. Comparing odds of overweight at age 7 years, boys in the lowest birthweight tertile with accelerated growth had odds of overweight at least as high as boys in the highest birthweight tertile without accelerated growth (odds ratio (OR)=2.50, 95% CI (1.71, 3.66) and OR=2.31 (1.61, 3.30), respectively). Across analyses, children in the highest birthweight tertile with accelerated growth experienced the highest odds of overweight, though accelerated growth was much less common among these children compared with children in the lowest birthweight tertile (14.5% vs 53.3%).

The second study, Jones-Smith *et al.*,⁴¹ examined BMI z-score at birth as a prenatal growth measure, any increase in linear BMI z-score between birth and 1 year as a measure of accelerated growth, and BMI and overweight at ages 4–6 years among Mexican children. They found significant interactions between BMI z-score at birth and increases in BMI z-score⁴¹ in multiple linear and logistic regression analyses (interaction terms were $b = -0.15$, $(-0.25, -0.05)$ and OR=0.64, (0.44, 0.90) respectively). A larger increase in BMI z-score in the presence of low or normal BMI z-score at birth, but not high BMI z-score at birth, was associated with higher odds of overweight at 4 to 6 years (low BMI z-score: OR 3.58, 95% CI (1.68, 7.44); normal BMI z-score: OR 2.23, 95% CI (1.12, 4.46); high BMI z-score: OR 1.41, 95% CI (0.63, 3.16)). Among children with low and normal BMI z-scores at birth, accelerated growth was associated with higher risks of overweight at ages 4–6 years compared with children with low and normal BMI z-scores at birth, respectively; however, among children with high BMI z-scores at birth, accelerated growth did not result in higher risks of overweight compared with children with high BMI z-scores at birth who did not experience rapid growth. Thus, accelerated growth in the first year of life conferred additional risk of overweight among children small or normal size at birth, but not among children who were large at birth. This result differs from other studies included in this review that found children born large who experience rapid growth typically have a greater risk of obesity compared with children born large who do not experience rapid growth.^{36,40,45,46} The conflicting finding reported by Jones-Smith *et al.*⁴¹ may be due to the definition of large size at birth: it used z-score to define size at birth, while others used size for gestational age or tertiles of birthweight. However, children in the Jones-Smith 2007 cohort who were large at birth (regardless of whether they experienced accelerated growth) still had the highest risk of overweight.

The third study, Gishti *et al.*, assessed birthweight for gestational age categories (SGA: <10 th percentile, AGA: ≥ 10 th–90th percentile, LGA: >90 th percentile), change in weight s.d. >0.67 from birth to 2 years (catch-up, catch-down and normal growth), and measured BMI, fat mass index, lean mass index and android/gynoid fat ratio at 6 years among children in the Netherlands.³⁶ Interaction terms between gestational age categories and change in weight categories were significantly associated with fat mass and android/gynoid fat ratio; no details are provided. In stratified analyses, SGA children with catch-up growth had significantly higher measures of android/gynoid fat ratio, reflecting central body fat distribution in the abdomen and hip regions, compared with AGA children with normal growth; they had values of android/gynoid fat ratio similar to AGA children with catch-up growth and to LGA children with normal growth (SGA: 0.22 (0.09, 0.35); AGA: 0.26 (0.18, 0.34); LGA: 0.21 (0.04, 0.38)). SGA children with catch-up growth resembled AGA children with normal

Table 3. Summary of Results: Associations between Prenatal Growth, Postnatal Growth, and Indicators of Obesity^a

	BMI or BMI z-score	Odds of overweight or obesity	Waist circumference	Fat mass or fat mass index	Relative fat mass	% body fat or fat mass	Central to total fat ratio ^b	Waist to height ratio	Waist-to-hip ratio or android/gynoid fat ratio	Sum of skinfold thickness ^c	Subscapular to triceps skinfold thickness ratio ^d	Ratio of preperitoneal to subcutaneous distance ^e	Central fat mass ^f	Peripheral fat mass ^f	Study
Prenatal growth															
BW	+	NR ^h +	+	NS	+										(33) (42) (46)
	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	(38)
	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	(31)
	+Boys only	+Boys only	+Boys only	NS	+Boys only	-Girls only	-	-	-	-	-	-	-	-	(35) (39) (34)
	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	(48)
	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	(32) (40)
BMI	+	+	+	+	+	+	NS	+	NR	NS	NR	NR	NR	NR	(45) (41)
Size for gestational age	NR	+	+	NR	NR	NR	NS	+	NR	NR	NR	NR	NR	NR	(37) (43)
IUGR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	(47) (36) (44)
Postnatal growth															
Δ weight	+	+	+	+	+	+	+	+	NR	NS	NR	NR	NR	NR	(45) (46) (34) (31) (33) ⁱ (39) (35) (40) ⁱ (48) (32) (47) (36) (38) (43) (44) (37) (41) (42)
	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	(45)
	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	(46)
	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	(34)
	+	+	+	+	+	+	+	+	NR	NS	NR	NR	NR	NR	(31)
	+	+	+	+	+	+	+	+	NR	+	NR	NR	NR	NR	(33) ⁱ (39)
	+	+	+	+	+	+	+	+	NR	+	NR	NR	NR	NR	(40) ⁱ (48)
	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	(32) (47)
	+	+	+	+	+	+	+	+	NR	NR	NR	NR	NR	NR	(36) (38) (43) (44) (37) (41) (42)
	+	NR	NS	NS	NS	NS	NS	NS	NR	NS	NR	NR	NR	NR	(31) ^j (32)
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Table 3. (Continued)

	BMI or BMI z-score	Odds of overweight or obesity	Waist circumference or obesity	Fat mass index	Fat-free mass or fat mass	% body fat or fat mass	Relative fat mass	Central to total fat ratio ^b	Waist to height ratio	Waist-to-hip ratio or android/gynoid fat ratio	Sum of skinfold thickness ^c	Subscapular to triceps skinfold thickness ratio ^d	Ratio of preperitoneal to subcutaneous distance ^e	Central fat mass	Peripheral fat mass ^f	Study	
BW × rapid growth	LBW: +, NS ABW: +, NS ^{9g} +P=0.01 ^{ah}		LBW: +, NS ABW: + ^{ah}													(38)	
BW × BMI IUGR × Rapid growth ^{ah}		P=0.78 non-IUGR: + Obesity: IUGR: NS; non-IUGR: NS ⁹ⁱ OW: P=0.42 Obesity: P=0.33 P=0.92															(40) (42) (44)
WFL × WFL																	(48)

Abbreviations: AGA, appropriate for gestational age; BMI, body mass index; BW, birthweight; CUG, catch-up growth; FM, fat mass; FMI, fat mass index (kg/m³); FFM, fat-free mass; FFMI, fat-free mass index (kg/m²); g, grams; IUGR, intrauterine growth restricted; kg, kilograms; LGA, large for gestational age; mos, months; OW, overweight; RG, rapid growth; s.d., standard deviation; SFT, skinfold thickness; SGA, small for gestational age; STR, subscapular to triceps skinfold thickness ratio; PP/SC-distance, preperitoneal to subcutaneous distance; WC, waist circumference; WFL, weight-for-length; WHR, waist-to-hip ratio; wks, weeks; Δ, change. ^aThis table presents a summary of results from all articles included in this review that assessed the independent effect of prenatal growth, while accounting for the effect of postnatal growth, the independent effect of postnatal growth, while accounting for the effect of prenatal growth, and the interaction between prenatal growth and postnatal growth. Symbols indicate the sign and significance of associations with each obesity-related outcome: significant positive association (+), significant negative association (-), and nonsignificant association (NS). Blanks cells indicate the association was not assessed. ^bThe central fat ratio refers to a ratio of the sum of suprailiacal and subscapular SFT to the sum of triceps and biceps SFT. ^cThe sum of four SFT refers to the sum of biceps, triceps, suprailiacal and subscapular skinfold thicknesses. ^dThe subscapular to triceps skinfold thickness ratio (STR) is used to assess regional differences in subcutaneous fat distribution, with a higher ratio representing increased central fat deposition. ^eThe ratio of preperitoneal to subcutaneous distance (PP/SC-distance) refers to the ratio between preperitoneal fat thickness and subcutaneous thickness, with a higher ratio representing increased a more adverse distribution of abdominal fat. ^fCentral fat mass refers to the sum of suprailiacal and subscapular SFT. Peripheral fat mass refers to the sum of triceps and biceps SFT. ^gResults were not significant for quartile 2 of postnatal growth with any outcomes but were positive for quartiles 3 and 4 with all outcomes. ^hThis association became nonsignificant after adjusted for breastfeeding and weight at 1 year. ⁱThese studies used multiple postnatal growth measurements. ^jSign and/or P-value of interaction term not reported. ^kIncrease in weight s.d. score >0.67. ^lWithin each BW tertile, CUG not significantly associated with adverse abdominal fat distribution (compared with 2nd tertile BW, normal growth): 1st tertile BW -4.40 (-15.89, 8.76); 2nd tertile BW -12.10 (-25.40, 3.56); 3rd tertile BW -4.78 (-36.17, 42.05), P trend = 0.02. ^mSGA with CUG no significant difference in BMI compared with AGA with normal growth -0.2 (-0.4, -0.1); LGA with continued growth had significantly increased BMI (0.8 (0.4, 1.1)). ⁿSGA with CUG no significant difference in WC compared with AGA with normal growth (-0.3 (-0.9, 0.3)); LGA with continued growth significantly increased WC compared with AGA with normal growth (1.9 (1.1, 2.8)). ^oSGA with CUG no significant difference in FMI compared with AGA with normal growth (-0.1 (-0.3, 0.05)); LGA with continued growth significantly increased FMI compared with AGA with normal growth (0.2 (0.01, 0.50)). ^pSGA with CUG no significant difference in % body fat compared with AGA with normal growth (-0.2 (-0.9, 0.4)); LGA with continued growth no significant difference in % body fat compared with AGA with normal growth (0.5 (-0.4, 1.5)). ^qSGA with CUG no significant difference in WHR compared with AGA with normal growth (0.005 (-0.001, 0.011)). ^rSGA with CUG significantly increased S:T ratio compared with AGA with normal growth (0.003 (-0.001, 0.007)); AGA with continued growth no significant difference in WHR compared to AGA with normal growth (-0.2 (-0.4, 0.01)). ^sP > 0.29 for all tests of interaction. ^t6 wks: b = -6.18 × 10⁵, P = 0.76; 6 mos: b = 0.00, P = 0.13. ^u-7.38 × 10⁻⁶, P = 0.059; postnatal weight gain has a larger effect on fat distribution for lower BW compared with higher BW. ^vSGA with CUG no significant difference in BMI compared with AGA with normal growth (0.00 (-0.11, 0.11)); AGA with CUG and LGA with CUG had significantly increased BMI (AGA: 0.55 (0.45, 0.59); LGA: 1.11 (0.64, 1.58)). ^wSGA with CUG had nonsignificant increase in FMI compared with AGA with normal growth (0.09 (-0.02, 0.20)); AGA with CUG and LGA with CUG had significantly increased FMI (AGA: 0.31 (0.24, 0.38); LGA: 0.73 (0.26, 1.19)). ^xSGA with CUG had nonsignificant decrease in FFMI compared with AGA with normal growth (-0.04 (-0.11, 0.02)); AGA with CUG and LGA with CUG had significant decreased FFMI (AGA: -0.16 (-0.20, -0.12); LGA: -0.30 (-0.56, -0.03)). ^ySGA with CUG had significant increase in android/gynoid fat ratio compared with AGA with normal growth (0.22 (0.09, 0.35)); AGA with CUG had significant increase in android/gynoid fat ratio (0.26 (0.18, 0.34)) while LGA with CUG had a nonsignificant increase in android/gynoid fat ratio (0.44 (-0.11, 0.99)). ^zComparison group = AGA with normal growth: SGA with CUG significantly lower BMI s.d. score (-0.23 (-0.39, -0.07)); AGA with CUG significantly higher BMI (0.44 (0.37, 0.52)) LGA with continued growth significantly higher BMI (1.06 (0.77, 1.36)). ^{aa}Comparison group = AGA with normal growth: SGA with CUG no significant difference in BMI compared with AGA with normal growth (0.00 (-0.26, -3.22, 2.70)); LGA with continued growth significantly higher SFT (3.50 (2.02, 4.97)); LGA with continued growth significantly higher SFT (8.95 (3.93, 13.97)). ^{ab}Comparison group = AGA with normal growth: SGA with CUG no significant difference (-0.11 (-1.52, 1.30)); AGA with CUG significantly higher central FM (1.95 (1.25, 2.76)); LGA with continued growth significantly higher central FM (4.42 (2.02, 6.81)). ^{ac}Comparison group = AGA with normal growth: SGA with CUG no significant difference (-0.15 (-2.11, 1.80)); AGA with CUG significantly higher peripheral FM (1.47 (0.51, 2.44)); LGA with continued growth significantly higher peripheral FM (4.55 (1.24, 7.87)). ^{ad}Interaction term: b = -0.15 (-0.25, -0.05), P < 0.01. ^{ae}Interaction term: OR = 0.63 (0.44, 0.90), P < 0.05. ^{af}No significant differences in mean BMI were observed for LBW with rapid growth (mean = 17.4, s.d. = 3.0). ^{ag}No significant differences in mean WC were observed for LBW with rapid growth (mean = 57.8, s.d. = 5.6) compared with LBW without rapid growth (mean = 54.1, s.d. = 3.3), compared with ABW without rapid growth (mean = 57.8, s.d. = 5.3) or compared with ABW with accelerated growth by BW tertile: low 2.50 (1.71, 3.66); med 3.98 (2.62, 6.05); high 4.97 (3.16, 7.83); odds of OW (ORs, 95% CI) for boys with accelerated growth by BW tertile: low 1.12 (0.73, 1.72); med 1.96 (1.22, 3.15); high 3.32 (1.85, 5.95). ^{ah}Increase in weight-for-age z-score > 0.67. ^{ai}Odds OW IUGR with RG: 2.69 (0.22, 32.51); odds obesity non-IUGR with RG: 1.63 (1.10, 2.43); odds obesity IUGR with RG: 2.69 (0.22, 32.51); odds obesity non-IUGR with RG: 1.79 (0.97, 3.30).

growth in BMI, fat mass index and lean mass index but had body fat more centrally distributed.

Figure 2 depicts odds of overweight and obesity among children aged 2–7 years according to prenatal and postnatal growth patterns among the three studies that presented odds of overweight or obesity by size at birth and speed of growth (see Hui *et al.*,⁴⁰ Jones-Smith⁴¹ and Taal *et al.*⁴⁷). Odds of overweight and obesity were highest for children born heavier who also grew rapidly. Children of normal birthweight who grew rapidly were at higher risk of overweight or obesity than children born heavier who then grew more slowly but had lower risks than children born heavier who then grew rapidly. Children born light (though not necessarily low birthweight) who experienced rapid growth had higher odds of overweight and obesity compared with children born light who experienced slow or normal growth; they also experienced higher odds of overweight and obesity compared with normal birthweight children with slow or normal growth; their risks approximated those of normal birthweight children with rapid growth.

Among the 15 studies that did not find significant interactions between prenatal and postnatal growth, four studies found no significantly higher risk of obesity in the presence of rapid postnatal growth among children or adolescents born SGA, IUGR, LBW stunted or LBW wasted.^{38,43,44,47} Monteiro *et al.*⁴⁴ reported that rapid growth was not significantly associated with overweight or obesity risks among adolescents (ages 14–16 years) who had experienced IUGR (overweight prevalence ratio (PR)=2.46 (0.70, 8.65); obesity: PR=2.69 (0.22, 32.51)), but it was positively associated with obesity among adolescents who had not experienced IUGR (PR=1.63 (1.10, 2.43)). This result warrants further study, since infants who experience IUGR have higher mortality than those who do not,⁵² which is a potential competing risk to obesity, potentially biasing results towards the null. Kramer *et al.*⁴³ compared SGA children who did and did not experience catch-up growth with AGA children who experienced normal growth and found no significant differences at age 11.5 years in BMI, percentage body fat, fat mass index, waist circumference or waist-to-hip ratio.⁴³ However, SGA children with catch-up growth resembled AGA children on almost all measures and had healthier measures on all outcomes than SGA children without catch-up growth, who remained significantly lighter, shorter, and thinner

than AGA children. Similarly, Taal *et al.*⁴⁷ found no difference between SGA children with catch-up growth and AGA children with normal growth in terms of odds of overweight, obesity or skinfold thickness measured at ages 2–4 years: for each group (SGA, AGA and LGA) catch-up growth resulted in higher fat mass and BMI. However, odds of obesity were similar among SGA children with catch-up growth and AGA children with normal or catch-down growth. Goncalves *et al.*³⁸ compared the association between rapid postnatal growth and BMI and waist circumference at age 8 years between children with appropriate BW (ABW) and children with low birthweight who were either stunted (16.6%) or wasted (22.1%) based on the Ponderal Index. LBW children with rapid growth had similar measures of BMI and waist circumference to ABW children with and without rapid growth. These results entail that postnatal growth may be less problematic for children or adolescents who are growth-restricted *in utero* than for others.

DISCUSSION

This review evaluated evidence of the independent and combined effects of prenatal growth and postnatal growth for childhood obesity. Results indicate that, accounting for subsequent postnatal growth patterns, prenatal growth is generally positively associated with obesity, though there are some inconsistencies when assessing adiposity and some differences between boys and girls. Accounting for prenatal growth, postnatal growth is positively associated with obesity and adiposity across the birthweight distribution. Of the 18 studies that have examined whether the implications of postnatal growth for obesity differ by size at birth, 15 reported that this association is similar for children/adolescents of all sizes at birth. However, three studies found that postnatal growth increased risks of obesity as much or more for infants at the lower end of the birthweight distribution as for infants born at a normal size or large size.

Across studies, children and adolescents who were born heavy and experienced rapid postnatal growth had the highest odds of obesity, highest BMIs and indicators of adiposity. Children and adolescents born in the normal range of birthweight who experienced rapid postnatal growth were also at higher risk of obesity-related outcomes, with risk approximating those of high birthweight children and adolescents who experienced slow

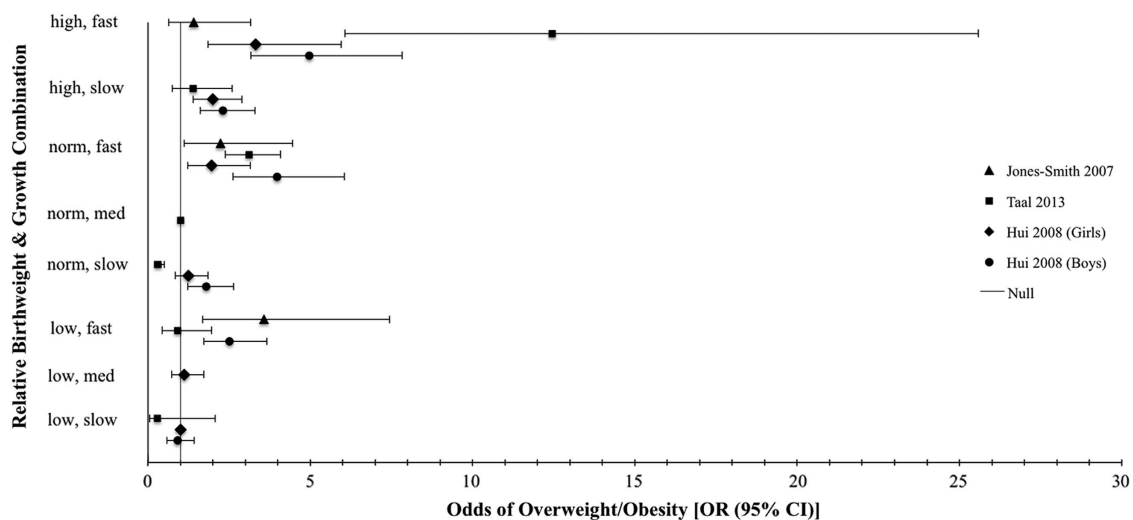


Figure 2. Odds of overweight and obesity among children aged 2–7 according to their relative birthweight and postnatal growth strata. Results are from three studies reporting odds ratios and 95% confidence intervals for overweight or obesity for each combination of birthweight, categorized as low, normal (norm) or high, and postnatal growth categorized as slow, medium (med) or fast. These strata reflect the relative sizes at birth for each report’s study population and do not correspond with clinical definitions of low birthweight (< 2500 g) or high birthweight (>4000 g). ▲ Jones-Smith 2007. ■ Taal 2013. ◆ Hui 2008 (Girls). ● Hui 2008 (Boys). — Null.

postnatal growth. Children and adolescents born lighter who experienced rapid growth may also be at higher risk of some obesity-related outcomes, although there is some evidence that those who are growth-restricted *in utero* may benefit from rapid postnatal growth. More research is needed to confirm the patterns indicated by previous studies to assess the balance between the risks and benefits of rapid growth, and to refine recommendations. These studies highlighted that it is important to monitor growth from birth to 2 years of age, as this is an important developmental period for all children. A priority for research will be to examine the implications of specific prenatal growth patterns and of specific types and timing of postnatal growth patterns; it will be important to examine patterns from across the distribution of prenatal and postnatal growth.

An important question is which groups should be compared to understand the implications of prenatal and postnatal growth for health: should low-birthweight infants who grow rapidly be compared with (1) low-birthweight infants who do not grow rapidly, (2) average-birthweight infants with average postnatal growth or (3) average-birthweight infants who grow rapidly? An informative approach is to make comparisons with ideal prenatal and postnatal growth to get an overall sense of risk (i.e. reference category being those with normal birthweight and normal postnatal growth). Another approach would be to make comparisons with a group that differs only on postnatal growth in order to determine whether rapid postnatal growth is beneficial or harmful given the growth pattern experienced before birth. There is currently no standard practice, and studies have used each of these approaches along with proxy measures for both prenatal and postnatal growth (such as birthweight and change in weight between two time points) that do not fully capture the dynamic patterns of infant growth. A first step to addressing this challenge will be to systematically assess each of these comparisons, preferably within the same population. This step will identify which groups are different from each other and where are the differences, when we hold other conditions constant. It will also elucidate which components of prenatal and postnatal growth are being captured with various measures and which are relevant for subsequent growth. A novel modeling method, SuperImposition by Translation and Rotation (SITAR), has been proposed as a way to independently examine factors that may influence growth parameters, such as size and velocity, in different ways.^{53,54} Latent class growth modeling is one method used recently to distinguish groups of children with biologically distinct growth trajectories and estimate the effects of prenatal and postnatal exposures on the odds of membership in a specific growth trajectory group.⁵⁵ Future research could benefit from utilizing these advanced methods of studying early life growth patterns.

Among the studies that found similar associations between postnatal growth and obesity for all infants, some features may entail that the findings are not definitive. One study had very few low-birthweight children,³² three did not report on the proportion of participants in their studies born low birthweight^{45–47} and one included no low-birthweight children.⁴³ For six of the studies, sample sizes of the subgroups are not shown, and the numbers of participants who were born small and then grew rapidly may be small.^{32,44–48} Samples with few participants born low-birthweight, as well as stratification of participants resulting in small subgroups, can result in an inability to detect significant differences across groups. Another study assessed effect modification only among adolescents born low-birthweight;³⁵ the inclusion of only those born low-birthweight precludes assessment of whether there is a different association between postnatal growth and obesity for those born low-birthweight compared with those born larger. Future studies could be designed and powered to be generalizable across the birthweight distribution.

Our study and the literature it described have limitations. Comparison of results between the studies is complicated by differences across them in measures of prenatal growth, postnatal growth and obesity-related outcomes. The periods of postnatal growth varied in duration and age, as did age at outcome assessment and reference standards used for overweight and obesity. Standard definitions and measures of prenatal and postnatal growth have not been uniformly adopted, nor have measures of obesity and adiposity in childhood and adolescence.

While studies have used measures such as weight at birth and BMI z-score at birth as measures of prenatal growth, these proxies are neither equivalent to each other nor accurately capture prenatal growth. A child of the same birthweight can have reached that birthweight through very different growth patterns, including unimpaired intrauterine growth or asymmetrical or symmetrical intrauterine growth retardation—all of which follow different growth patterns and may have different implications for subsequent metabolic conditions. To better understand the relationships between prenatal growth and obesity, future studies should consistently measure prenatal growth by gestational age, as has also been previously recommended.^{22,56} Studies that use other proxies, such as birthweight, should be very clear about what such a measure can and cannot tell us about prenatal growth. Lack of standard terminology is a major concern: several studies used the same terms for different measures of postnatal growth and others used different terms for the same measure. For example, of studies reporting on 'rapid growth', one used the term to describe an increase in weight-for-age z-score greater than 0.67 units,⁴⁴ while the other used it to refer to the highest quintiles of weight gained per month;⁴⁶ a third study used the same cutoff but named it 'accelerated growth'.⁴⁰ Adoption of standard definitions and measures of prenatal growth, postnatal growth and obesity-related indicators using a standard reference population will help solidify our understanding of these topics. For example, though it is often difficult to discern between physiology and pathology, the term catch-up growth should be used for a physiological condition of temporary overgrowth, as catch-up growth takes the child back onto his or her original growth centile trajectory; in contrast, the term rapid growth should be limited to denote an abnormal condition of overgrowth.

Research is needed to better understand associations between prenatal growth, postnatal growth and obesity, and whether growth patterns have different implications for girls and boys, as one of the studies suggested. It will also be important to determine which aspects of body size and body composition may be affected by prenatal or postnatal growth, and whether rapid postnatal growth is linked with a general increase in body size or a selective increase in some aspect of body composition. The mechanisms by which early life growth patterns are associated with subsequent adiposity require more research, including an examination of the links between environmental and behavioral factors and early life growth patterns, such as sleep, activity levels and family resources.⁵⁷

Based on the findings summarized here, we can conclude that early life growth patterns have implications for future disease risk, with prenatal and postnatal periods each contributing to the development of later obesity. Prenatal and postnatal growth patterns could be used to construct an early life risk profile to identify children who may benefit from early intervention to reduce the risk of chronic disease later in life.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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APPENDIX

Table A1. Correlations between prenatal growth and postnatal growth^a

	Change in weight (kg)	Rate of weight gain (g/month)	Weight z-score	Change in weight s.d. score	Change in weight z-score tertiles	Change in weight for length z-score quartiles	Change in weight z-score > 0.67	Change in weight-for-age z-score > 0.67	Change in BMI z-score	Δ kg/m ² /per year	Study
BW (g or kg)		Stettler <i>et al.</i> ⁴⁶		– ^b			– ^c				(33) (38) (42) (46)
BW s.d. score; z-score			– ^e	– ^f NR – ^g							(31) (34) (35) (39)
BW for length z-score BW (tertiles)					– ⁱ	+ ^h	NR				(48) (32) (40)
BW (quartiles) BMI z-score	NR								NR – ^d		(45) (37) (41)
Size for gestational age							NR – ^j	NR			(36) (43) (47)
IUGR								NR			(44)

Abbreviations: BMI, body mass index; BW, birthweight; g, grams; IUGR, intrauterine growth restricted; kg, kilograms; r, correlation coefficient; s.d., standard deviation; Δ, change. ^aThis table summarizes results from articles that reported the correlation between prenatal growth and postnatal growth. Symbols indicate the following correlations between prenatal growth and postnatal growth were measured: positive (+), negative (–), relationship not reported (NR). Blank cells indicate the correlation was not assessed. ^b $r = -0.48$, $P < 0.001$. ^cSignificantly higher proportion of LBW children experienced rapid weight gain than children of appropriate BW (LBW stunted: 51.9%, LBW wasted: 75.0%, ABW: 28.0%, $P < 0.001$). ^dResults not shown. ^e $r = -0.53$. ^fLower BW SDS inversely correlated with change in weight SDS: 0–3 wks $r = -0.36^*$; 3–6 wks: $r = -0.19^{**}$; 6–12 wks: $r = -0.12$; 3–6 mos: $r = -0.16^{**}$; 6–12 mos: $r = -0.07$; $*P < 0.001$, $**P < 0.05$. ^g0–6 wks: $r = -0.35$; 0–6 mos: $r = -0.56$, $P < 0.001$. ^hChildren with higher 6 mos WFL z-score more likely to have higher birth WFL z-score, $P = 0.06$. ⁱLowest BW tertile had highest percentage of fast growers 0–3 months (53.3% vs 14.5% for highest BW tertile). ^jSGA children had greater change in weight SDS from birth to 3 months than AGA (1.12 (0.96, 1.29), $P < 0.001$); LGA children experienced a decrease in weight SDS from birth to 3 months compared with AGA children (–1.25 (–1.41, –1.10), $P < 0.001$).