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## **Review Article**

## Health impact of catch-up growth in low-birth weight infants: systematic review, evidence appraisal, and meta-analysis

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#### Abstract

This study aimed to systematically review and appraise evidence on the short-term (e.g. morbidity, mortality) and long-term (obesity and non-communicable diseases, NCDs) health consequences of catch-up growth (vs. no catch-up growth) in individuals with a history of low birth weight (LBW). We searched MEDLINE, EMBASE, Global Health, CINAHL plus, Cochrane Library, ProQuest Dissertations and Thesis and reference lists. Study quality was assessed using the risk of bias assessment tool from the Agency for Health Care Research and Quality, and the evidence base was assessed using the GRADE tool. Eight studies in seven cohorts (two from high-income countries, five from low-middle-income countries) met the inclusion criteria for short-term (mean age: 13.4 months) and/or longer-term (mean age: 11.1 years) health outcomes of catch-up growth, which had occurred by 24 or 59 months. Of five studies on short-term health outcomes, three found positive associations between weight catch-up growth and body mass and/or glucose metabolism; one suggested reduced risk of hospitalisation and mortality with catch-up growth. Three studies on longer-term health outcomes found catch-up growth were associated with higher body mass, BMI or cholesterol. GRADE assessment suggested that evidence quantity and quality were low. Catch-up growth following LBW may have benefits for the individual with LBW in the short term, and may have adverse population health impacts in the long-term, but the evidence is limited. Future cohort studies could address the question of the consequences of catch-up growth following LBW more convincingly, with a view to informing future prevention of obesity and NCDs. © 2016 John Wiley & Sons Ltd

Keywords: obesity, NCDs, infant feeding, catch-up growth, low birthweight.

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## Introduction

Low birth weight (LBW), defined by the WHO as a birth weight <2500 g (UNICEF, WHO 2004), is common, particularly in low-middle-income countries (LMICs). It is clear that LBW typically leads to poor health outcomes. Conservative estimates of LBW prevalence made by UNICEF and the WHO in 2004 suggested that at least 16% of births globally were LBW, with around 96% of these in LMICs (UNICEF, WHO 2004).

Accelerated postnatal 'catch-up' growth (in length, weight or both) is a common compensatory mechanism for LBW, which occurs typically in the first 24 months of postnatal life (Crowther *et al.* 1998; Jaquet *et al.* 2005).

It is believed that catch-up growth is beneficial for the individual in the short term (Victora *et al.* 2001), but may create public health problems in the long term because it may be associated with metabolic disturbances, which increase the risk of some non-communicable diseases (NCDs) and obesity (Kramer *et al.* 2014; Jain & Singhal 2012). It is believed that early catch-up growth, before around the age of 2 years, is beneficial for long-term health outcomes, but catch-up growth that occurs later than around 2 years increases risk of later obesity and NCDs (Victora *et al.* 2008), but this evidence has not focused on individuals with LBW and has not been subject to systematic review and evidence appraisal. The extent to which catch-up growth might influence short-term and long-term outcomes following LBW is therefore a major public-health-nutrition question, of particular importance for obesity and NCD prevention in LMICs.

Whether, and to what extent, catch-up growth following LBW in early life should be considered in future policy responses to the obesity and NCD crisis depends on the quantity, quality and consistency of the evidence relating catch-up growth following LBW to short-term and long-term health outcomes. No previous systematic review has considered differences in health outcomes following LBW in those with catch-up growth vs. those without catch-up growth. One review (Nobili et al. 2008), generated from a literature search in a single database, compared the effect of catch-up growth in LBW vs. non-LBW individuals, but did not compare outcomes for individuals born LBW with catch-up growth vs. those without catch-up growth. A recent analysis of data from five birth cohorts in LMICs, not focused specifically on those born LBW, suggested that catch-up growth after 2 years of age would increase later risk of obesity and NCDs (Adair et al. 2013).

The primary aim of this study was therefore to examine the impact of catch-up growth (vs. no catch-up growth) on health outcomes in those born LBW. A secondary aim was to critique the available evidence, identifying gaps and weaknesses, so that future studies might permit a more confident assessment of the impact of catch-up growth following LBW, as part of a more evidence-informed global approach to NCD and obesity prevention in the future.

## Methods

# Eligibility criteria: studies; study participants; exposures and outcomes

All study designs were eligible for inclusion in this review so long as they provided data for infants and children where catch-up growth occurred prior to 59 months, with a history of LBW as defined by the WHO (birth weight < 2500 g), only studies with participants who had a history of LBW as defined by WHO were included. Definitions of catch-up growth vary between studies, and no international standard has been established. Study eligibility was therefore not limited by the definition of catch-up growth used, and studies were included so long as catch-up growth was defined (including definitions based on Weight-for-age; Height-for-age; Weight-for-height).

The following outcomes were considered: direct measures of adiposity and proxies for adiposity; blood pressure; fasting blood glucose; impaired glucose tolerance; elevated glycosylated haemoglobin (HbA1c); insulin and insulin resistance; total blood cholesterol, triglycerides, lipoprotein levels (low density lipoprotein – LDL, high-density lipoprotein – HDL) and cardiometabolic risk scores, which included any or all of the previous indicators. Eligible measures of cardiovascular events were angina pectoris, stroke, myocardial infarct and mortality. Risk of diabetes type 2 was also included.

#### Search methods for identification of studies

We searched the following electronic databases on 6 August 2014: MEDLINE (1946 to July week 4 2014); EMBASE (1974 to 2014 week 31); Global Health (1910 to 2014 week 30); CINAHL plus (1983 to August 2014); Cochrane Library (up to issue 7 of 12 July 2014); ProQuest Dissertations & Theses (1980 to August 2014). The journal Bulletin of the WHO was searched in Pubmed Central (1948 to 1st June 2014), and a hand search of the WHO South-East Asian Journal of Public Health and the publication lists of birth cohorts listed at http://www.birthcohorts.net/ was performed. In addition, we examined reference lists and citations of relevant studies. A search for new studies that had cited eligible studies was carried out in November 2015, but produced no additional eligible studies. Keywords were searched

### Key messages

- Some evidence supports the view that early life catch-up growth (compared to no catch-up growth) following LBW
  is beneficial in the short-term, but harmful in the long-term
- The evidence base is small (8 eligible studies), relatively low quality, and not entirely consistent
- Making a strong case for the avoidance of catch-up growth as a target of NCD and obesity prevention strategy would not be evidence-based at present

as subject headings indexed in databases and as free-text terms. Booleans were used to refine the search. The search strategy for Medline is given later (Fig. S1). Controlled vocabulary and search syntax were modified as appropriate when searching other databases. Only studies in the English language were included.

#### Data collection, management and analysis

#### Selection of studies

AM and AC screened and cross-checked titles and abstracts independently to identify potentially relevant studies based on the previous criteria. Full-text reports of potentially relevant studies were assessed for eligibility independently by two reviewers (AM, JJR). Discrepancies were resolved by discussion and where needed, RMB arbitrated. A list of excluded studies was generated and reasons for exclusion recorded.

#### Data extraction and management

We used a standardised protocol for extracting relevant information from the studies. Data extraction was performed independently by two reviewers (AM and JJR) who resolved any differences by discussion.

#### Quality assessment of included studies

Quality of included studies was assessed independently by AM and JJR, cross-checked and discussed to resolve disagreement where required. We used the 10-item risk of bias-assessment tool from the Agency for HealthCare Research and Quality (Viswanathan *et al.* 2013) to assess study quality formally.

#### Assessment of publication bias

If the number of included studies allowed ( $\geq 10$  studies), we aimed to assess reporting bias by using a funnel plot.

#### Data synthesis and quality assessment of evidence

Available data were not suitable for meta-analysis, with the exception of two studies that examined weight-forage and height-for-age catch-up associations with fasting insulin (see later). Weighted mean differences of insulin levels between children with and without catch-up growth were combined using random-effect models to account for unobserved variables. Review manager 5.3 was used for data synthesis (Review Manager (RevMan) [Computer program] 2014). Where studies were considered insufficiently similar to each other to be combined in a meta-analysis, results were described by timing of outcome (short term up to the age of 5 years; longer term after 5 years). Estimates of effects were summarised in the GRADE Evidence Profile (Brozek *et al.* 2008) along with the quality rating of the evidence.

Where studies did not report the statistical significance of the group difference (between those with a history of LBW with catch-up growth vs. those with a history of LBW without catch-up growth), and where data were available, data were re-analysed to determine significance of a group difference using inverse variance and random-effect models.

### Results

#### Search outcomes

The searching and screening process is summarised in Fig. 1. The literature search yielded 881 records, of which 283 were duplicates. Titles and abstracts of 598 records were screened, resulting in 98 records for fulltext screening (86 papers and 12 abstracts). Independent screening and cross-checking (AM, JJR) identified eight eligible studies for inclusion; 90 records did not meet the inclusion criteria and thus were excluded. Reasons for exclusion are listed in Fig. 1.

#### Characteristics of included studies

Included studies are summarised in Table 1a and b for short-term and longer-term outcomes, respectively.

#### General study characteristics

Of the eight studies (seven cohorts), five were prospective, and three were cross-sectional. Evidence was available from two studies in high-income countries and six (from five cohorts) from LMICs.

#### Population

The total number of children studied was 535 (short-term health outcomes; Table 1a) and 553 (longer-term health outcomes; Table 1b). LBW was defined by

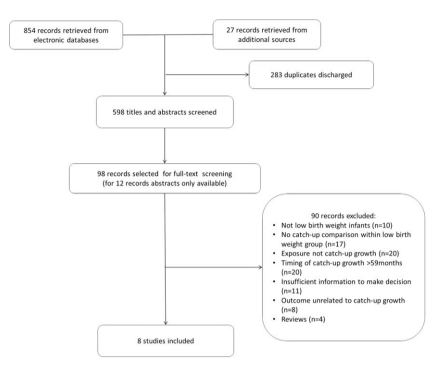


Fig. I Literature Search: Study Flow Diagram.

individual studies as: birth weight or length < 10th percentile of a sex and gestational age specific reference (Horta *et al.* 2003; Han *et al.* 2010; Rustogi *et al.* 2013; Victora *et al.* 2001); weight < 5th percentile for gestational age (Soto *et al.* 2003; Rustogi *et al.* 2013); weight and/or length < 2SD below means for gestational age (Tenhola *et al.* 2000) and birthweight < 2500 g (Khandelwal *et al.* 2014; Mai *et al.* 2005). In all of the eligible studies, participants met the WHO definition of LBW. Attrition rates of participants ranged from 16% to 86% with a median of 27%. Two studies did not report how many children were lost to follow-up (Han *et al.* 2010; Rustogi *et al.* 2013).

#### Exposure

Dichotomous definitions of catch-up growth (comparing those who 'caught up' with those who did not) were used, but with different cut-offs to distinguish between those who caught up and those who did not: weight and/or height gain of  $\geq 0.67$  z-scores (Khandelwal *et al.* 2014; Rustogi *et al.* 2013; Soto *et al.* 2003; Victora *et al.* 2001; Horta *et al.* 2003), or weight or height z-score increase from birth-follow-up of  $\geq 2$  (Tenhola *et al.* 2000) or >0(Han *et al.* 2010). All included studies reported outcomes related to weight catch-up growth, while three also reported on height/length catch-up growth (Han *et al.* 2010; Rustogi *et al.* 2013; Soto *et al.* 2003), and one provided additional data on weight-for-height catch-up growth (Rustogi *et al.* 2013). Seven studies reported on catch-up growth up to the age of 24 months and three studies included children who caught up after 24 months.

#### Comparison

All but three studies reported the impact of catch-up growth on markers of obesity or NCD risk compared with children who did not catch-up. Three studies provided data on the impact of change in weight *z*-scores between two time points on obesity, NCD risk or risk or markers of NCDs (Horta *et al.* 2003; Khandelwal *et al.* 2014; Mai *et al.* 2005).

#### Outcomes

Of the nine eligible studies, five tested for associations between catch-up growth and early health outcomes (Han *et al.* 2010; Khandelwal *et al.* 2014; Rustogi *et al.* 2013; Soto *et al.* 2003; Victora *et al.* 2001; early outcomes defined here and pre-specified as aged < 5 years), while four tested for associations between catch-up growth and later health outcomes (Horta *et al.* 2003; Mai *et al.* 2005; Tenhola *et al.* 2000; Victora *et al.* 2001; later defined

Table I.	(a) Characten	stics and sh	iort-term health ou	Table I. (a) Charactenstics and short-term health outcomes of included studies							
Study ID	Study characteristics	istics				Participant characteristics			Exposure- catch-up growth		
	Study Design	Study location	Recruitment setting	Exclusion criteria	Attrition rates	Low BW/SGA definition	Mean BW	Term / preterm	Definition	Type	Timing
<b>Han et al.</b> 2010	Cross- sectional	Peking, China	Third Hospital, Peking University	not singletom, gestational age <33 wks. non SGA, 1-min Apgar score <7, 5 min Apgar score <10, intratterine infectors, ongenital malformations, major neonatal problems, breastfed <3 months, mothers with diabetes,	29%	below <10th percentile of the sex speci- fic distribution for gestational age using birth weight standards of Chinese	1996.59 g (353.15)	gestational age of > 33 weeks (mean 36.46 SD2.38wk)	The change of weight Z- score during the 3 months > 0 Z-score was defined as catch-up growth	weight	3 то
Khandelwal et al. 2014	Prospective cohort study	India	not reported	gestational diabetes, chronic hypertention birth weight <1500 g, breast feeding not possible, requirement of intravenous fluids, ambiotiss, oxygene or CUS stuy for more than 24 h at birth, mujor congenital malformations, stigmata of intrauterine infections, genetic syndromes or chromosomal anomal ics and	50%	BW <2500 g	2175 ± 180 g, z score -2.67 ± 0.49	term: gestational age between 37 and 42 weeks	Changes in weight z score between birth and the follow up visits difference in z score > 0.67 in weight for age (AWAZ)	Weight WAZ	1.4 mo 3 mo 7.2mo 1.4 mo 3 mo
Rustogi et al. 2013	Cross – sectional study	India	not reported	reatence more than 40 km from the study site not reported	not reported	weight or length <10thpercentile	not reported	term	gain in weight length SDS or both of $>0.67$	weight length weight /	12-18 mo 12-18 mo 12-18
<b>Soto et al.</b> 2003	Cross - sectional	Chile	neonatal units of Hospital San Borja Arriara'n and Hospital So'tero	significant medical, neurological, or genetic conditions, on unusual diets or were taking any medication that could interfere with growth or appetite	Not reported	birth weight <5th percentile for gestational age, using Chilean birth weight standards	2.1 SDS ± 0.1	term: Gestational age 37-41 wks	weight/lengths gain, between zero and 1 yr, greater than 0.67 SDS	weight height	1y 1y
<b>Victora <i>et al.</i></b> 2001	Prospective cohort study	Pelotas, Brazil	det Kt o households	notreported	15%	BW <10th centile of weight for gestational age of the Williams curve	not reported	not reported for SGA	weight change in zscores > =0.66 from bi rth to 20 months	weight	20 mo
(b) <b>Horta <i>et al.</i></b> 2003	prospective cohort study	Pelotas, Brazil	5 maternity hospitals	not reported	86%	< 10th centile for gestational age and sex, according to the reference developed by Williams et al	not reported	not reported for SGA	Changes in weight z score between birth and the follow up visits	weight	20 mo 42 mo 20 mo
<b>Mai et al.</b> 2005	prospective cohort	Sweden	hospitals	not reported	16%	VLBW <1500 g	not reported	not reported	changes of SDS in weight between postmenstrual age of 40 wk and fol lowup time points (6 months, 10	weight	-2 шо 6 mo 18mo
Tenhola <i>et</i> <i>al.</i> 2000 Victora <i>et al.</i> 2001	Prospective cohort study Prospective cohort study	Finland Pelotas, Brazil	Kuopio University Hospital households	Metabolic Disease not reported	25% 15%	birth weight and/or length and/or ponderal index <2 SD score below the respective mean for the gestational age BW <10th centile of weight for gestational age of the Williams curve	median 2452 g (2367, 2537) not reported	term not reported for SGA	weight or height increase > = 2.8D score between birth and 5y weight change in zscores > =0.66 from birth to	weight height weight	5y 5y 20 mo
									20 months		

BW: birth weight, SGA: small-for-gestational age, mo: months, y: year, B: unstandardized regression coefficient, B; standardized regression coefficient, OR: odds ratio, SD: standard deviation, CI: Confidence interval

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	Outcome measure	Time point	No catch-up group		Catch-up group		p-value of difference	Confounders
<b>Han</b> et al. 2010	Fasting glucose (mmol/L)	3 mo	n = 12	Mean (SD): 4.18 (0.58)	n = 32	Mean (SD): 4.32 (0.64)	0.528	none
	Insulin sensi tivi ty (HOMA)		n = 12	4.15 (2.96)	$n = 32 \ 2.11 \ (1.06)$	n = 32 2.11 (1.06)	0.356	
Khandelwal <i>et al.</i> 2014	FM%	7.2 mo	n = 33	$\beta = 2.91, 95\%$ CI - 0.88 to 6.70			0.13	gender, current age and
		7.2 mo	n = 33	$\beta = 5.00, 95\%$ CI 0.67 to 9.33			0.03	current length
		7.2 mo	n = 33	$\beta = 5.42, 95\%$ CI 1.43 to 9.43			0.01	
		7.2 mo	n = 14	Mean (SD) = 12.8 (7.6)	n = 6	Mean $(SD) = 21.4 (7.5)$	0.06	
		7.2 mo	n = 14	Mean (SD) = 12.8 (7.6)	n = 13	Mean $(SD) = 18.5(7.5)$	Not reported	
Rustogi et al. 2013	fasting insul in (uIU/ml)	12-18mo	n = 32	Mean $(SD) = 3.0 (2.5)$	n = 18	Mean(SD) = 7.3(9.2)	0.01	none
	fasting insul in (uIU/ml)	12-18mo	n = 25	Mean $(SD) = 3.2 (2.2)$	n = 25	Mean (SD) = 5.9(8.3)	0.2	none
	fasting insul in (uIU/ml)	12-18mo	n = 20	Mean $(SD) = 2.8 (1.9)$	n = 30	Mean(SD) = 5.8(7.6)	0.06	none
Soto et al. 2003	BMI (kg/m2)	1y	n = 22	Mean $(SD) = 15.9 (0.2)$	n = 63	Mean(SD) = 17.2(0.2)	<0.001	none
	fasting insul in (pmol/L)		n = 22	Mean (SD): 14.9 (2.3)	n = 63	Mean(SD) = 32.6(4.6)	<0.001	
	Insulin sensi tivi ty AUC		n = 22	Mean (SD)= 2215.4(461.6)	n = 63	Mean(SD) = 2302.6	0.4	
	(pmol/minxL)							
	Insulin sensi tivi ty (1st phase		n = 22	Mean $(SD) = 303.5 (91.2)$	n = 63	Mean(SD) = 298.8(46.4)	0.82	
	insulin release, pmol/L)							
	BMI (kg/m 2)		n = 41	Mean $(SD) = 16.8(0.2)$	n = 44	Mean (SD) = 16.8(0.2)	1	
	fasting insul in (pmol/L)	1y	n = 41	Mean $(SD) = 20.9 (2.1)$	n = 44	Mean(SD) = 34.6(6.5)	<0.001	
	Insulin sensi tivi ty AUC		n = 41	Mean $(SD) = 1767.6(199)$	n = 44	Mean(SD) = 2790.8(400.9)	<0.001	
	(pmol/minxL)							
	Insulin sensi tivi ty (1st phase insulin release - pmol/L)		n = 41	Mean (SD) = 223.4 (27.3)	n = 44	Mean (SD) = 374.8 (76.4)	<0.001	
Victora et al. 2001	Al l-cause Hospital	30 mo	n = 25	Proportion of chi ldren 16.00%	n = 304	5.60%	Not reported	family income, maternal
	admissions							schooling, age
	Diarrhoea - hospi tal		n = 25	0.00%	n = 304	0.00%	Not reported	
	dutiliosi Otos							
	Lower respi ratory Infections – hospital admissions		n = 25	4.00%	n = 304	2.30%	Not reported	none
(q)								
Horta et al. 2003	systolic blood pressure	15y	total $n = 101$	B = -0.49; 95% CI 4.80 to 3.82 B = 1.86, 95%CI -2.91 to 6.64			not reported	family income, duration of breast feeding, gender,
	diastol ic blood pressure			B = -0.01; 95% CI - 4.21 to 4.20				maternal height, and
				B = -0.32, 95% CI -4.98 to 4.34				maternal smoking during
								pregnancy
Mai et al. 2005	BMI (kg/m2)	12 y	n = 74	correlation: $\rho = 0.34$			<0.01	None
		12y	n = 74	correlation: $\rho = 0.24$			<0.05	none
Tenhola <i>et al.</i> 2000	high cholesterol levels (> 4.8 mM)	12y	total $n = 34$	OR 0.3, 95% CI 0.1 to 1.9		OR 1.0	0.3	none
	high cholesterol levels (> 4.8 mM)	12y	total $n = 35$	OR 13.8, 95% CI 2.0 to 97.5		OR 1.0	0.009	
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here and pre-specified as aged  $\geq$  5 years); one of the eligible studies included both short-term and longer-term outcomes (Victora *et al.* 2001). The following NCD risk factors were assessed: BMI (Mai *et al.* 2005; Soto *et al.* 2003; percentage fat (Khandelwal *et al.* 2014); glucose metabolism (Han *et al.* 2010; Rustogi *et al.* 2013; Soto *et al.* 2003); blood pressure (Horta *et al.* 2003); plasma cholesterol (Tenhola *et al.* 2000) and hospital admissions and mortality (Victora *et al.* 2001).

#### Quality appraisal of included studies

Overall, the quality across all included studies was low. Only two studies met five (i.e. low risk of bias) out of the 10 quality criteria; the remaining studies met less than five quality criteria. Attrition bias (applicable for cohort studies only) and selective reporting bias were not addressed by included studies, and bias due to confounding was only rarely addressed.

#### Selection bias

None of the included studies were at risk of selection bias. Children with or without catch-up growth were from the same cohort and thus quality item 2 was not applicable (differing recruitment strategy for individuals).

#### Detection bias

All studies failed to provide adequate details on whether the assessor was blinded to the exposure or outcome, and thus, the studies were judged to be of 'unclear' risk of bias. Six out of nine studies used valid and reliable measures of exposure and outcome and thus were of low risk of bias. However, three studies were judged as 'unclear' as insufficient information was reported (Horta *et al.* 2003; Rustogi *et al.* 2013; Victora *et al.* 2001).

#### Attrition bias

Attrition bias was not applicable in the longitudinal studies which used cross-sectional analyses (Han *et al.* 2010; Rustogi *et al.* 2013; Soto *et al.* 2003). The remaining prospective studies showed no differences in followup time between comparison groups. However, three of the prospective studies did not assess the impact of attrition, which was high (>20%), with potential to bias the outcome (Horta *et al.* 2003; Khandelwal *et al.* 2014; Tenhola *et al.* 2000). Thus, these studies were at high risk of attrition bias. A further two studies did not assess the impact of attrition; however, their attrition rates were low and so less likely to bias the results (Mai *et al.* 2005; Victora *et al.* 2001). Therefore, the risk of attrition bias was low.

#### Selective reporting bias

The majority of studies did not refer to a published study protocol which would allow assessment of whether all predetermined outcome measures were reported. Thus for these studies the risk of selection bias was judged to be 'unclear' (Han *et al.* 2010; Horta *et al.* 2003; Mai *et al.* 2005; Rustogi *et al.* 2013; Victora *et al.* 2001). For three studies, it was possible to determine that relevant outcomes were not reported (Khandelwal *et al.* 2014; Soto *et al.* 2003; Tenhola *et al.* 2000); thus, the risk of selective reporting was judged to be high. Assessment of missing adverse events or harms was not applicable to all included studies.

#### Bias due to confounding

One study took known confounding factors into account when analysing the association between catchup growth and non-communicable disease risk factors and so was judged to be of low risk of confounding bias (Horta *et al.* 2003). The remaining studies did not account for confounders and were therefore considered to be at high risk of bias.

#### Synthesis of evidence

Most studies showed a high level of heterogeneity in terms of study design, length of follow-up, definition of the catch-up growth, timing of catch-up growth and outcomes assessed. Therefore, a quantitative synthesis of the evidence in a meta-analysis was not suitable except for one outcome measure. The evidence is described largely narratively by timing of outcome assessment below.

# Short-term outcomes of catch-up growth in low birth weight children

Of the studies that provided data on short-term outcomes, all referred to weight catch-up growth, only two studies (Rustogi *et al.* 2013; Soto *et al.* 2003) assessed the association of length/height catch-up growth on short-term health. Findings for weight and/or length catch-up growth can be found in Table 1a (by study) and 2a (by outcome). Reported short-term outcomes were hospital admission, body mass and glucose metabolism up to the age of 30 months; the mean age at outcome measurement was 13.4 months.

One study suggested that catch-up growth was associated with reduced risk of hospitalisation: hospitalisation (all-cause) was significantly lower in children with catch-up growth (n = 304) compared with children without (n = 25; Victora *et al.* 2001). Two studies found significantly higher fat mass by 5.7% (95% CI 0.0 to 11.4%; *n* = 27; Khandelwal *et al.* 2014) and BMI by  $1.30 \text{ kg/m}^2$  (95% CI 1.20 to  $1.40 \text{ kg/m}^2$ , n = 85; Soto et al. 2003) in children with catch-up growth compared with children without catch-up growth at 3 and 12 months, respectively. Three studies assessed the association between catch-up growth and glucose metabolism (fasting glucose or insulin or insulin sensitivity; Han et al. 2010; Rustogi et al. 2013; Soto et al. 2003). One study found no association between catch-up growth and fasting glucose (Han et al. 2010). Metaanalysis of the other two studies indicated higher fasting insulin levels of 2.54 uIU/mL (95% CI 2.33 to 2.76 uIU/mL, P < 0.001,  $I^2 = 0\%$ ) in children with weight catch-up growth (n = 50) compared with the no weight catch-up growth group (n = 54). Individual study findings on the association between height catch-up growth and fasting insulin were inconclusive. However, pooled mean differences showed higher fasting insulin levels of 2.00 uIU/mL (95% CI 1.70 to 2.29 uIU/mL,  $P < 0.001, I^2 = 0\%$ ) in children with height/length catch-up growth. Insulin sensitivity was more impaired in children without weight and/or height catch-up growth compared with children that showed weight and/or height catch-up growth at 3 months (Rustogi et al. 2013) and 12 months (Soto et al. 2003, Table 2a).

## Longer-term outcomes of catch-up growth in low birth weight children

Longer-term outcomes were available for weight catchup growth from all studies and for height catch-up growth by one study (Tenhola *et al.* 2000). Reported longer-term outcomes between 5 and 15 years (mean age 10.2 years) were mortality, body mass index, blood pressure and cholesterol levels. Findings are summarised for each study in Table 1b and by outcome in Table 3b.

Based on one single study (Victora et al. 2001), mortality by the age of 5 years was (non-significantly) lower in children with catch-up growth compared with those with no catch-up growth. BMI at age 12 years was significantly correlated with changes in weight z-scores between birth and 6 months and between birth and 18 months (n = 74). The correlation coefficients were 0.34 and 0.24, respectively (Mai et al. 2005). There was no evidence of a significant association between catch-up growth and diastolic blood pressure at 15 years in one study (n = 101; Horta *et al.* 2003). Children with height (not weight) catch-up growth (n=21)had a 13.8 fold (95% CI 2.0 to 97.5) increased risk of high total cholesterol levels of >4.8 mM/L at 12 years compared with children without catch-up growth (n = 35; Tenhola et al. 2000).

#### Quality and consistency of evidence

The GRADE evidence profiles for short- and longterm outcomes are summarised in Table 2a and 3b, respectively. The quality of evidence was very low for the outcomes percent body fat, BMI, glucose levels, insulin levels, insulin sensitivity, systolic and diastolic blood pressure, risk of high cholesterol levels for height catch-up growth and low for hospital admissions and mortality. The reason for the grades of very low to low quality was because evidence was available from predominantly low-quality observational studies only. Evidence inconsistency could not be adequately assessed because for almost all outcomes only one or two studies were eligible.

### Discussion

#### Main study findings and implications

The present study found a relatively small body of evidence of low to very low quality according to AHRQ and GRADE methodology, which addressed the question of the impact of catch-up growth (vs. no catch-up

Quality assessment	nent						No of patients	nts	Effect		Quality
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Catch-up growth	No catch-up growth	Relative (95% CI)	Absolute	
Percentage fat	Percentage fat mass – weight catch-up at 3 months (follow-up 5.8 months)	-up at 3 months (	follow-up 5.8 mor	aths)							
1	observational	serious <sup>1</sup>	no serious	no serious	serious <sup>2</sup>	none	13	14		MD 5.7% higher	000⊕
	study		inconsistency	indirectness							VERY LOW
Body Mass Inc	Body Mass Index - weight catch-up at 12 months	p at 12 months									
1	observational	serious <sup>3</sup>	no serious	no serious	no serious	none	63	22		$MD 1.30 \text{ kg/m}^2$	000⊕
	study		inconsistency	indirectness	imprecision					higher (1.20 to 1.40 higher)	VERY LOW
Body Mass Inc	Body Mass Index - height catch-up at 12 months	) at 12 months								0	
1	observational	serious <sup>1</sup>	no serious	no serious	no serious	none	44	41		$MD 0.00 \text{ kg/m}^2$	000⊕
	study		inconsistency	indirectness	imprecision					higher (-0.09 to 0.09)	VERY LOW
Fasting glucose	Fasting glucose - weight catch-up at 3 months	at 3 months									
1	observational	serious <sup>4</sup>	no serious	no serious	no serious	none	32	12		MD 0.14 mmol/L	000⊕
	study		inconsistency	indirectness	imprecision					higher	VERY LOW
Insulin sensitiv	Insulin sensitivity levels – weight catch-up $3$ months (HOMA) and $12$ months (AUC)	ttch-up 3 months	(HOMA) and 12	months (AUC)							
2	observational	serious <sup>1</sup>	no serious	no serious	serious <sup>5</sup>	none	95	34		HOMA: MD 2.04	000⊕
	studies		inconsistency	indirectness						higher AUC: MD	VERY LOW
Fasting insulin	Fasting insulin levels – weight catch-up at 12–18 month	12-18 mo	onths							o'	
2	observational	serious <sup>3</sup>	no serious	no serious	no serious	none	50	54	I	Not pooled: mean	000⊕
	studies		inconsistency	indirectness	imprecision					ranged from 2.6 to	VERY LOW
	(cross-sectional)									4.3 uIU/mL higher	
Hospital admis	Hospital admission - weight catch-up at 20 months (follow-up mean 10 months)	np at 20 months (	follow-up mean 1	0 months)							
1	observational	no serious	no serious	no serious	no serious	none	304	25		10.4% lower	⊕000 L0W
	study	risk of bias	inconsistency	indirectness	imprecision						

<sup>1</sup>Studies did not account for attrition and confounding variables, there was evidence of selective outcome reporting.

<sup>2</sup>Wide confidence intervals indicate imprecision, The sample size was low.

<sup>3</sup>Study did not account for confounders and selective reporting of outcomes was evident. <sup>4</sup>Study did not account for confounding variables. <sup>5</sup>Low sample size in the comparison group is likely to add imprecision to the overall effect.

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Quality assessment	sessment						No of patients	nts	Effect		Quality
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Catch-up growth	No catch-up growth	Relative (95% CI)	Absolute	
Body Mas	Body Mass Index - weight catch-up at 6 and 18 months	atch-up at 6 an		(follow-up 10.5-11.5 years)	us)						
1	observational	serious <sup>1</sup>	no serious	serious <sup>2</sup>	no serious	none				Correlation 0.34	000⊕
	study		inconsistency		imprecision		74			to 0.24 higher	VERY LOW
Systolic bl	ood pressure - we	ight catch-up a	Systolic blood pressure - weight catch-up at 20 months (study 2) (follow-up mean 13.3 years)	am dn-wollo) (follow-up me	an 13.3 years)						
1	observational	serious <sup>3</sup>	no serious	serious <sup>2</sup>	serious <sup>5</sup>	none				B = 0.49  mmHG	000⊕
	study		inconsistency				101			lower (-4.80 to 3.82)	VERY LOW
Diastolic <b>k</b>	olood pressure - w	eight catch-up	Diastolic blood pressure - weight catch-up at 20 months (follow-up mean 13.3 years)	v-up mean 13.3 y	ears)						
1	observational	serious <sup>3</sup>	no serious	serious <sup>2</sup>	serious <sup>5</sup>	none				B = 0.01  mmHG	000⊕
	study		inconsistency				101			lower (-4.21 to 4.2)	VERY LOW
Systolic bl	ood pressure – we	sight catch-up a	Systolic blood pressure – weight catch-up at 42 months (study 2)	2)							
1	observational	serious <sup>3</sup>	no serious	serious <sup>2</sup>	serious <sup>5</sup>	none				B = 1.86  mmHG	000⊕
	study		inconsistency				101			higher (-2.91 to 6.64)	VERY LOW
Diastolic h	olood pressure – w	eight catch-up	Diastolic blood pressure – weight catch-up at 42 months (follow-up mean 12.5 years)	v-up mean 12.5 y	(ears)						
1	observational	serious <sup>3</sup>	no serious	serious <sup>2</sup>	serious <sup>5</sup>	none				0.32 lower (4.98 lower	000⊕
	studies		inconsistency				101			to 4.34 higher)	VERY LOW
Cholester	ol levels – weight (	atch-up at 59 n	Cholesterol levels – weight catch-up at 59 months (follow-up mean 7 years)	tean 7 years)							
1	observational	serious <sup>3</sup>	no serious	no serious	no serious	none	21/55	34/55	OR 0.3	291 fewer per 1000	000⊕
	studies		inconsistency	indirectness	imprecision		(38.2%)	(61.8%)	(0.1 to 1.9)	(from 479 more to	VERY LOW
			\$		4		~	~	~	136 more)	
Cholester	ol levels – height c	atch-up at 59 n	Cholesterol levels – height catch-up at 59 months (follow-up mean 7 years)	ean 7 years)							
1	observational	serious <sup>3</sup>	no serious	no serious	serious <sup>5</sup>	strong association <sup>6</sup>	20/55	35/55	OR 13.8	324 more per 1000	000⊕
	studies		inconsistency	indirectness		reduced effect for $BP > > 1$ or	(36.4%)	(63.6%)	(2 to 97.5)	(from 141 more to 358 more)	VERY LOW
						RR << 1					
Mortality	- weight catch-up	at 59 months (1	Mortality – weight catch-up at 59 months (follow-up mean 3.3 years)	years)							
1	observational	no serious	no serious	no serious	no serious	none	304	25	ļ	75% lower (3 vs.	⊕000 LOW
	studies	risk of bias	inconsistency	indirectness	imprecision					13 less per 1000)	
<sup>1</sup> Study dic <sup>2</sup> Study ass <sup>3</sup> Studies di <sup>4</sup> Study ass	<sup>1</sup> Study did not account for confounding variables. <sup>2</sup> Study assessed effect of change in weight <i>z</i> -score over <sup>3</sup> Studies did not account for confounding and attrition. <sup>4</sup> Study assessed the effect of change in weight <i>z</i> -score ri	onfounding vai unge in weight , confounding a change in weig	<sup>1</sup> Study did not account for confounding variables. <sup>2</sup> Study assessed effect of change in weight z-score over time rather than effect of catch-up vs. no catch-up. <sup>3</sup> Studies did not account for confounding and attrition. <sup>4</sup> Study assessed the effect of change in weight z-score rather than effect of weight catch-up vs. no weight catch-up.	ther than effect a	of catch-up vs. n tht catch-up vs. r	o catch-up. 10 weight catch-up.					
Wide cor Study rep	<sup>o</sup> Wide confidence intervals indicati <sup>6</sup> Study reported a large effect size.	ndicate imprec 2t size.	"Wide confidence intervals indicate imprecision, the sample size was small. <sup>6</sup> Study reported a large effect size.	e was small.							

Table 3. (b) GRADE evidence profile for long-term outcomes of catch-up growth

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growth) in LBW infants on short-term and longer-term health outcomes. No previous systematic review addressed this research question. For some of the studies, the main research questions were not the same as the research questions addressed by the present review. In addition, for studies conceived, conducted, and/or reported prior to the recent widespread use of AHRQ and GRADE methodology, low study quality was likely due in part to the age of the studies and lack of awareness of the methodology.

Consistency of the evidence is hard to assess because, for almost all of the outcomes, only single studies were available. With limited quantity and quality of evidence, and uncertainty over the consistency of the evidence, it cannot be concluded that catch-up growth following LBW increases risk of adverse cardiometabolic health in later life. Long-term-outcome data, in adults, were missing.

#### Limitations of the review

Meta-analysis of the studies identified in the present review was limited to one outcome and only two studies because of substantial heterogeneity between studies and lack of data on the same outcome measure. Publication bias could not be assessed formally because the number of eligible studies was too small. It may be of note that included studies reported both significant and non-significant associations of catch-up growth vs. no catch-up growth on health outcomes of relatively small participant number. Thus, the presence of publication bias on the grounds of effect sizes and study impact is less likely. We had planned subgroup-analyses, e.g. examining differences by age, exposure characteristics such as being LBW as a result of being born too small for gestational age or appropriate for gestational age, gender, setting, study design and sensitivity analyses (synthesising all of the available evidence and then only those studies deemed to have low risk of bias), but the small number of eligible studies, and their heterogeneity, precluded such analyses. This review focused solely on research published in English language, and thus, potentially relevant studies published in other languages might have been missed. Translating records into English language was not feasible for this review.

## Limitations of the evidence base and implications for future research

The research question asked by the present review is an important one for global public health nutrition, regardless of whether or not it can be answered with any great confidence at present. In order to answer it with evidence of higher quality, future research should address the issues summarised in Table 4, namely, (i) many of the eligible studies made no reference to study power; (ii) many failed to take into account confounders, despite potentially important differences between those with catch-up growth vs. no catch up growth (e.g. greater prevalence or severity of morbidity

Table 4. Summary of research suggestions for population, exposure, comparison, outcomes and data analysis

Population	Exposure	Comparison	Outcomes	Data analysis
More research on low birth weight infants needed	Standardised definitions of length catch-up growth and weight catch up growth;	Need for more research specifically comparing those with low birth weight and catch up	Need for more evidence on a range of outcomes, but particularly adult health outcomes	Multivariate regression analysis taking potential confounding variables into account
More focus on subgroups within the low birth weight population (e.g. SGA and AGA)	More emphasis on trajectories of catch up;	growth vs. LBW with no catch up growth		Consideration of attrition and missing outcome data in data analysis
Increased sample size to increase statistical power Reporting of reasons of attrition (e.g. mortality, drop out, moving away)	More emphasis on growth and anthropometric end points (e.g. catch up growth to height or length within the healthy range vs. stunting)			

in the latter); (iii) many studies did not account for attrition; (iv) substantial heterogeneity in the definitions of catch-up make it difficult to understand what exposure actually matters; and (v) there was substantial heterogeneity inherent in the exposure. The LBW definition included individuals of widely varying birth weight, timing of catch-up growth will have varied, and includes both those born too early and those born too small – an important distinction (Lapillone & Griffin 2013) that was made by some studies (Table 1) but not all.

A large number of ineligible studies compared catchup growth of LBW children with growth of children born at or above 2500 g (Fig. 1). Studies that were excluded because they did not meet the comparison group criterion might have suitable data available to answer the research question asked by the present study. Some studies that did not meet our inclusion criteria for other reasons can also provide useful evidence. Kramer et al. (2014) did not compare formally between those who showed catch-up growth vs. those who did not, but noted that those who caught-up had slightly higher adiposity than those who did not. In one large study from the USA, Hemachandra et al. (2007) treated catch-up growth as a continuous exposure variable, with no comparison between those who showed catch-up growth vs. those who did not (so was ineligible here), but reported that those with higher gains in weight z-score in infancy and early childhood had significantly increased risk of high-blood pressure at age 7 years.

There is a need for a clearer understanding of the nature and timing of the exposure of catch-up, more evidence on the short-term and long-term impacts of catch-up growth vs. no catch-up growth in LBW infants, and whether the consequences of catch-up vary between children with a history of LBW vs. those without. Researchers with access to existing (or planned cohorts) might consider this research question in future in order to address the evidence gaps identified by this review. Specific questions, such as the importance of the precise timing or rate of catch-up growth, the relative importance of length vs. weight catch-up growth, whether health outcomes of catch-up growth differ for those born too early vs. those born too small, and the mechanisms that relate catch-up growth to later health outcomes, could not be answered.

## Conclusions

In summary, the present study has found some evidence that catch-up growth in those born LBW is beneficial relative to no catch-up in the short-term. The longer-term population health impact of catch up growth (vs. no catch up growth) in those born LBW is less clear. Major weaknesses and gaps in the evidence, combined with the importance of the issue of catch-up growth to global population health, demonstrate that further studies, or secondary analyses of available data, are required urgently.

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## **Conflicts of interest**

The authors declare that they have no conflict of interest.

## Contributions

Initial searching and screening: AM and AC; Full text screening: AM, JJR, RMB; Drafting of manuscript: JJR and AM; Redrafting of manuscript for critical content: all authors.

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