There is now compelling evidence that growth patterns in early life are associated with risk of the metabolic syndrome in adulthood, although the relative importance of prenatal vs. postnatal growth for such associations remains controversial. Body composition may play a key role in the ‘programming’ of such diseases, through itself being programmed by early growth, and perhaps also by being a mediator of the programming process. Early studies reporting positive associations between birth weight and adult BMI suggested a tendency for large babies to become obese adults. Such findings appeared contradictory to the many studies linking low birth weight with increased risk of the metabolic syndrome. Recent studies now indicate that birth weight is strongly predictive of later lean mass, and has a much weaker association with later fatness. Studies that link low birth weight with a more central adipose distribution in later life remain controversial, and require confirmation using more sophisticated methodologies. Findings for infant growth rate appear population-specific, with infant weight gain predicting subsequent lean mass in developing countries, but predicting subsequent fat mass and obesity in industrialised populations. Further studies are required on this issue, to ensure that appropriate public health policies are recommended for countries across the range of economic development. Although the links between early growth and later disease risk implicate early-life nutrition, either in utero or during infancy, few prospective studies have explored the influence of early diet on later body composition. Many studies have associated breast-feeding with a reduced prevalence of obesity categorised by BMI; however, the few studies directly evaluating childhood fatness provide little support for this hypothesis. Recent advances in the ability to measure body composition during the infant period offer a major opportunity to improve the understanding of the nutritional programming of body composition and its contribution, or lack thereof, to subsequent disease risk.

Body composition: Nutritional programming: Birth weight: Infant growth rate

In the last two decades a large volume of evidence has emerged associating growth patterns during early life with subsequent risk of the metabolic syndrome, implicating early-life nutrition as the underlying mechanism (Barker et al. 2002; Wells, 2007). The importance of early development for later health outcomes is reflected in the concept of programming, i.e. the notion that during early ontogeny the developing organism passes through ‘critical windows’ of sensitivity or plasticity, during which environmental factors generate long-lasting variability in phenotype (Lucas, 1991).

Whilst the relative importance of fetal vs. postnatal growth for later disease risk remains controversial, it is no longer doubted that experience during early life is predictive of a variety of health outcomes, including body size, body composition and the risk of diseases such as hypertension, stroke, type 2 diabetes, obesity and CVD (Barker et al. 2002). Such work is supported by complementary research, much of it involving experimental interventions, in a variety of animal species (McMillen & Robinson, 2005).

The majority of studies associating early development with later disease have used birth weight as an index of fetal growth, with low birth weight being shown to be predictive of increased subsequent disease risk. In many cases the strongest association between birth weight and later outcome is found when a statistical adjustment for adult size is incorporated (Barker et al. 2002). This issue has more than one implication. First, it has been argued (Lucas et al. 1999) that such statistical models do not demonstrate that low birth weight is the key risk factor, rather that change in size between birth and adulthood is
most important. Thus, postnatal growth rate as well as fetal growth rate appears to be implicated in the disease process. Such a perspective does not discount the importance of fetal growth, since the worst outcome is often found in those babies born small who subsequently become large (Adair & Cole, 2002; Barker et al. 2002). However, low-birth-weight babies usually display the fastest rate of infant growth, while a faster trajectory of childhood growth has also been linked with increased disease risk (Barker et al. 2005).

The second issue raised by the statistical analyses is that adult BMI often makes an important contribution to the interaction between early growth patterns and subsequent disease risk. Adult obesity is now well established as a risk factor for the metabolic syndrome, and BMI is widely used as an index of obesity; however, across the entire range of body weight there is a correlation between BMI and lean mass as well as fatness. Furthermore, BMI is not very strongly correlated with the central abdominal fat, which is considered most harmful to health (Wells et al. 2007). The strong statistical contribution of BMI to analyses associating early growth with later outcomes does not, therefore, reveal which components, or components, of body composition are most important. Since epidemiological studies have typically used diseases as their primary outcome, there is a need to improve the understanding of how tissues and organs themselves are programmed. Early growth patterns might induce long-term effects on body composition, e.g. by impacting on hormonal axes that in turn regulate childhood growth. Alternatively, body composition variability might be a central component of early growth variability, and track from fetal life or infancy onwards.

Over the last decade an increasing number of studies have explored the programming of body composition in more detail. The primary aim here is to review such research, considering different periods of growth and their impact on total tissue masses and their regional distribution. A secondary aim is to consider the potential role of early-life nutrition in accounting for such associations. However, before reviewing this literature it is important to consider the difficulty of evaluating body composition in the epidemiological studies that are required to investigate such associations.

**Indices of body composition in epidemiological research**

The large sample sizes that are characteristic of epidemiological research generally preclude access to the most sophisticated body-composition techniques. Studies are therefore obliged to rely on simple techniques, which often only assess the outcome indirectly. Body composition comprises not only tissue mass but also aspects of size and shape. However, the ability to differentiate these variables becomes more difficult when indices of size and shape are themselves used to predict tissue mass.

The limitations of BMI as an index of adiposity are becoming increasingly appreciated. In children, for example, individuals of the same age and gender have a twofold range of body fat mass for a given BMI value (Wells 2000), as shown in Fig. 1(a), a scenario also present in those who are obese (Wells et al. 2006). Some children with high BMI are ‘stocky’ (high muscle mass) rather than fat, while other children within the normal BMI range nevertheless have high fat mass and relatively low muscle mass. BMI is therefore a poor outcome in epidemiological research investigating the possible programming of body composition and obesity. BMI also fails to reflect body shape and hence fat distribution, as shown in Fig. 1(b). Here, unpublished data for women from the UK National Sizing Survey (JCK Wells, P Treleaven and TJ Cole, unpublished results) illustrate how, after adjusting for hip girth, waist girth increases substantially with age within any given level of BMI.

Central adiposity has often been evaluated using the waist:hip ratio, or the triceps:subscapular skinfold thickness. These outcomes are likewise problematic in epidemiological analyses focusing on early growth. Differences between individuals in the waist:hip ratio may reflect physique (hip girth) as much as abdominal adiposity (waist girth). Fig. 1(c) plots waist circumference z-score v. hip circumference z-score in approximately 2000 young adult women from the UK National Sizing Survey (JCK Wells, P Treleaven and TJ Cole, unpublished results). A high waist:hip ratio is present in all those individuals with values lying above the regression line; however, many of those with such a high waist:hip ratio nevertheless have a low waist girth. Skinfold thickness ratios have likewise been shown to have poor statistical validity. The relationship between limb and torso skinfolds is not linear, and dividing one skinfold by another is a problematic way of assessing relative fat distribution (Wells & Victora, 2005). Similarly, percentage fat is a problematic way to express data for whole-body composition, since it reflects lean mass as well as fat and conceals absolute adiposity (Wells & Victora, 2005).

These findings emphasise the importance when investigating the early-life programming of body composition of using both improved methodologies for measuring body composition and more-appropriate statistical analyses.

**Fetal growth and later body composition**

Interest in the relationship between size at birth and later body composition is not derived only from the proposed importance of low birth weight in the programming process. For obvious practical reasons size at birth offers the earliest opportunity to assess growth rate and body composition in detail. Paradoxically, however, remarkably few studies have obtained data more sophisticated than weight and length at birth, while it is also now appreciated that birth weight may not play a causal role in associations between early experience and later outcome. Birth weight is best regarded as an outcome of convenience (Wilcox, 2001), and its value in indexing fetal growth is confounded by a lack of knowledge about genetic potential. As discussed later (see p. 426), researchers have adopted different approaches to address these problems, but birth weight remains an imperfect index of fetal growth.

A much-cited source of evidence concerning the programming of body composition during fetal life comprises
During the Second World War several populations were exposed to substantially-reduced energy intakes for specific time periods. Studies of the Dutch "hunger winter" (Stein et al. 2004) and the siege of Leningrad (Stanner & Yudkin, 2001) have shown that maternal undernutrition during pregnancy is associated with drastic reductions in the rate of conception, and with deficits in mean birth weight of approximately 300–500 g. Analysing the Dutch data according to the trimester of undernutrition suggests that the majority of this weight deficit can be attributed to famine exposure during the third trimester (Stein et al. 2004). Subsequent follow-up of the Dutch population that focused on adult males at the age of military conscription (Ravelli et al. 1976) has found between-group differences in the prevalence of obesity categorised according to BMI. Those individuals exposed in early pregnancy (first and second trimesters) were found to have an increased prevalence of adult obesity relative to non-exposed subjects, whereas those exposed in late pregnancy or early infancy have a reduced obesity prevalence. More recent follow-ups (Ravelli et al. 1999) have found similar results for waist circumference in women but not men. However, these data are difficult to interpret. First, the findings were not replicated in survivors of the Leningrad siege, who were found to show no association between prenatal exposure to famine and subsequent obesity risk (Stanner & Yudkin, 2001). Such negative findings have been attributed to the lack of catch-up growth in the Leningrad cohort, in contrast to the Dutch cohort (Stanner & Yudkin, 2001). Second, there is no need to invoke the concept of fetal programming of obesity to explain these results. An alternative interpretation posits selective conception according to maternal genotype or phenotype (maternal fatness), or selective survival according to fetal genotype or phenotype (offspring thriftiness). Thus, associations between maternal malnutrition, birth weight and later obesity do not necessarily implicate fetal growth patterns as the underlying mechanism, and may simply represent the greater capacity of initially-fatter women to conceive during harsh conditions.

A second and stronger source of evidence for the fetal programming of body composition derives from investigation of the effect of gestational age at birth. It is well established that preterm infants have low levels of body fat at birth (Rigo et al. 1998), a condition that can be attributed to the fact that fat deposition occurs largely during the follow-up of individuals exposed to malnutrition in utero. During the Second World War several populations were exposed to substantially-reduced energy intakes for specific time periods. Studies of the Dutch "hunger winter" (Stein et al. 2004) and the siege of Leningrad (Stanner & Yudkin, 2001) have shown that maternal undernutrition during pregnancy is associated with drastic reductions in the rate of conception, and with deficits in mean birth weight of approximately 300–500 g. Analysing the Dutch data according to the trimester of undernutrition suggests that the majority of this weight deficit can be attributed to famine exposure during the third trimester (Stein et al. 2004). Subsequent follow-up of the Dutch population that focused on adult males at the age of military conscription (Ravelli et al. 1976) has found between-group differences in the prevalence of obesity categorised according to BMI. Those individuals exposed in early pregnancy (first and second trimesters) were found to have an increased prevalence of adult obesity relative to non-exposed subjects, whereas those exposed in late pregnancy or early infancy have a reduced obesity prevalence. More recent follow-ups (Ravelli et al. 1999) have found similar results for waist circumference in women but not men. However, these data are difficult to interpret. First, the findings were not replicated in survivors of the Leningrad siege, who were found to show no association between prenatal exposure to famine and subsequent obesity risk (Stanner & Yudkin, 2001). Such negative findings have been attributed to the lack of catch-up growth in the Leningrad cohort, in contrast to the Dutch cohort (Stanner & Yudkin, 2001). Second, there is no need to invoke the concept of fetal programming of obesity to explain these results. An alternative interpretation posits selective conception according to maternal genotype or phenotype (maternal fatness), or selective survival according to fetal genotype or phenotype (offspring thriftiness). Thus, associations between maternal malnutrition, birth weight and later obesity do not necessarily implicate fetal growth patterns as the underlying mechanism, and may simply represent the greater capacity of initially-fatter women to conceive during harsh conditions.

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final trimester of pregnancy. However, by the time term age is reached preterm infants often continue to show low levels of body fat (Uthaya et al. 2005; although this outcome depends on early postnatal diet), a trait which is preserved into mid-childhood (Fewtrell et al. 2004). These findings are consistent with the hypothesis that the final trimester of pregnancy is an important period for the programming of fat mass. Despite their reduced total adipose tissue mass, MRI studies have suggested that by term age preterm infants have a more central adipose tissue distribution (Uthaya et al. 2005), with some indication that this trait persists subsequently (MS Fewtrell, A Lucas, TJ Cole and JCK Wells, unpublished results).

A third source of evidence for the fetal programming of body composition derives from studies of maternal metabolic control during pregnancy. Maternal type 2 diabetes and gestational diabetes are both associated with an increased risk of obesity in the offspring (Rodrigues et al. 1998; Whitaker et al. 1998). The influence of type 2 diabetes is more complex, and it might plausibly involve either a programming mechanism or genetic transmission (the thrifty genotype hypothesis). Recent studies of the Pima Indians (Dabalea et al. 2000; Dabalea & Pettit, 2001), a population characterised by unusually high rates of the disease, have elegantly differentiated between these pathways. Within mothers who at some stage develop type 2 diabetes the risk of the offspring developing the condition is much greater in those born after the diagnosis relative to those born before the diagnosis (Dabalea et al. 2000). The intrauterine diabetic environment is thus associated with an increased risk of obesity in both childhood and adulthood, independently of genetic factors and effects on birth weight (Dabalea & Pettit, 2001).

There is therefore some indication that adipose tissue mass and distribution may be programmed by experience during fetal life, independently of the contribution subsequently generated by postnatal growth pattern. These fetal effects may function relatively independently of birth weight, the importance of which remains controversial. A recent study (Hemachandra & Klebanoff, 2006) has shown, for example, that birth weight is reduced following intrauterine growth retardation during the third trimester of pregnancy, but is not reduced if growth retardation occurs during the first trimester of pregnancy. Thus, the ability of birth weight to indicate fetal growth experience is crude, and derives merely from the fact that since all embryos begin life at approximately similar size, birth weight must in some sense represent the magnitude of fetal growth. This proposition is confounded first by the lack of information about body composition, and second by the fact that fetal genotype influences fetal growth, such that birth weight cannot indicate the extent to which genetic growth potential has been achieved. One solution to this dilemma is to use ponderal index rather than birth weight, as a better index of nutritional status (for example, see Wells et al. 2005; Rogers et al. 2006). Another approach has been to categorise intrauterine growth retardation on the basis of disparity between birth weight and adult height (Leon et al. 1996). Such a strategy is ideal for studies of body composition development, but to the authors' knowledge this approach has not yet been adopted when considering body composition. Instead, the vast majority of studies have simply considered associations between birth weight and later BMI or body composition.

An extensive systematic review of factors associated with childhood obesity (Parsons et al. 1999) has identified numerous studies associating birth weight with later BMI, with each kilogram increase in birth weight typically increasing BMI by 0.5–0.7 kg/m$^2$. As BMI is the most widely-used categorisation for overweight and obesity, such studies were initially interpreted as indicating that fatter babies subsequently remain fatter. Such a conclusion would be consistent with studies associating maternal gestational diabetes with an increased risk of macrosomic offspring, who do indeed tend to remain obese (Wells, 2007). However, the scenario for diabetic mothers should not be taken to imply a linear association between birth weight and later fatness across the entire range of birth size. Some studies (Rogers & EURO-B LCS Study Group, 2003) have suggested ‘J’-shaped or ‘U’-shaped associations between birth weight and subsequent obesity risk categorised by BMI, implying a high prevalence of obesity in those of low or high birth weight. A smaller set of studies has explored associations between birth weight and subsequent regional distribution of fat. However, the findings were inconsistent and, like those mentioned earlier, mostly relied on simple anthropometry to assess the outcome.

Over the last decade, there has been a substantial rise in the number of studies exploring the relationship between birth weight and later body composition in greater detail. These studies have differed in the age range considered, the methods used to assess body composition and in the statistical approaches to analysing the data. Table 1 summarises such studies, addressing indices of lean mass, total body fatness and central adiposity. Despite using a variety of measurement techniques, these studies are fairly consistent, where appropriate data is available, in demonstrating significant associations between birth weight and subsequent lean mass. This association is broadly present across populations, reported, for example, in European and American studies as well as those from Brazil, Guatemala and India, and is apparent across the entire lifespan, the one exception being a study of infants born preterm (Fewtrell et al. 2004), possibly suggesting that preterm delivery disrupts the normal developmental trajectory of lean mass.

In contrast, the findings have been markedly less consistent for body fat and its distribution, with studies variously finding negative, positive or non-significant associations with birth weight (Table 1). This inconsistency appears attributable to a number of factors. First, methodological and statistical variability between studies is implicated (see p. 427, Table 1). Second, results appear to vary according to the time of follow-up. Third, two large studies from developing countries indicate a gender difference in such associations, with birth weight only associated with fat mass in females (Li et al. 2003; Sachdev et al. 2005). Similar findings have emerged from the Avon Longitudinal Study of Parents and Children cohort (Rogers et al. 2006).

Most such studies have focused on the entire spectrum of birth weight, and despite long-standing interest in low
birth weight as a predictor of late ill health, few studies have examined in detail the effects of low birth weight on later body composition. Those studies that have done so have confirmed the general association between birth weight and later lean mass but have shown weak or non-significant associations with later fatness (Hediger et al. 1998; Sachdev et al. 2005; Wells et al. 2005). One study (Kensara et al. 2005) has reported an increased percentage fat in low-birth-weight elderly adults after adjusting for adult BMI, consistent with several others reporting a general inverse association between birth weight and later fatness (Table 1). However, it remains

### Table 1. Associations of birth weight with body composition indices during childhood, adolescence, young adulthood and old age

<table>
<thead>
<tr>
<th>Reference</th>
<th>Population</th>
<th>n</th>
<th>Age (years)</th>
<th>Outcomes</th>
<th>Design</th>
<th>Lean</th>
<th>Fat</th>
<th>Central fat</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Childhood</strong></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hediger et al. (1998)</td>
<td>USA, M + F</td>
<td>4431</td>
<td>2–47 months</td>
<td>AA, SKF</td>
<td>R</td>
<td>+</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Okosun et al. (2000)</td>
<td>USA, M + F</td>
<td>2488</td>
<td>5–11</td>
<td>SKF</td>
<td>R</td>
<td>+</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>Mulligan et al. (2005)</td>
<td>UK, M + F</td>
<td>85</td>
<td>6–9</td>
<td>DXA, BP</td>
<td>R</td>
<td>–</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bavelkar et al. (1999)</td>
<td>India, M + F</td>
<td>477</td>
<td>8</td>
<td>BMI, WHR, SKF</td>
<td>R</td>
<td>+</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>Malina et al. (1996)</td>
<td>USA, M + F</td>
<td>237</td>
<td>7–12</td>
<td>BMI, SKF</td>
<td>PC</td>
<td>–</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Fewtrell et al. (2004)</td>
<td>UK, M + F, preterm</td>
<td>497</td>
<td>8–12</td>
<td>DXA</td>
<td>R, TT</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Wells et al. (2005)</td>
<td>Brazil, M</td>
<td>172</td>
<td>9–10</td>
<td>BMI, BIA</td>
<td>ANOVA, R</td>
<td>+</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Rogers et al. (2006)</td>
<td>UK, M + F</td>
<td>6086</td>
<td>9–10</td>
<td>DXA</td>
<td>R</td>
<td>+</td>
<td>+F</td>
<td>0</td>
</tr>
<tr>
<td>Singhal et al. (2003b)</td>
<td>UK, M + F</td>
<td>164</td>
<td>7–16</td>
<td>BIA, DXA, SKF</td>
<td>R</td>
<td>+</td>
<td>0</td>
<td></td>
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<tr>
<td><strong>Adolescence</strong></td>
<td></td>
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<tr>
<td>Labayen et al. (2006)</td>
<td>Spain, M + F</td>
<td>234</td>
<td>13–18</td>
<td>DXA, SKF</td>
<td>R</td>
<td>+</td>
<td>+</td>
<td>–M</td>
</tr>
<tr>
<td>Barker et al. (1997)</td>
<td>UK, M + F</td>
<td>216</td>
<td>14–16</td>
<td>BMI, WHR, SKF</td>
<td>R</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>Matthes et al. (1996)</td>
<td>UK, M + F</td>
<td>165</td>
<td>15–7</td>
<td>BMI, SKF</td>
<td>CC</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Frisrancho (2000)</td>
<td>USA, M + F</td>
<td>1993</td>
<td>15–17</td>
<td>BMI, SKF</td>
<td>R</td>
<td>–</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Kahn et al. (2000)</td>
<td>USA, M</td>
<td>192</td>
<td>17–22</td>
<td>BMI, WC, TA</td>
<td>R</td>
<td>+</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Euser et al. (2005)</td>
<td>The Netherlands, M + F</td>
<td>403</td>
<td>19</td>
<td>WH, BMI, SKF</td>
<td>R</td>
<td>+</td>
<td>0</td>
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<tr>
<td><strong>Young adulthood</strong></td>
<td></td>
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</tr>
<tr>
<td>Li et al. (2003)</td>
<td>Guatemala, M + F</td>
<td>267</td>
<td>21–27</td>
<td>SKF, WHR*</td>
<td>R</td>
<td>+</td>
<td>0</td>
<td>+F</td>
</tr>
<tr>
<td>Sachdev et al. (2005)</td>
<td>India, M + F</td>
<td>1526</td>
<td>26–32</td>
<td>BMI, SKF, WHR</td>
<td>PC</td>
<td>+</td>
<td>+F</td>
<td>–</td>
</tr>
<tr>
<td>Weyer et al. (2002)</td>
<td>USA Pima</td>
<td>272</td>
<td>25</td>
<td>DXA, UWW</td>
<td>R</td>
<td>+</td>
<td>0</td>
<td></td>
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<tr>
<td>Loos et al. (2001)</td>
<td>Belgium, M</td>
<td>229, twin pairs</td>
<td>18–34</td>
<td>BIA, SKF</td>
<td>R</td>
<td>+</td>
<td>–†</td>
<td>–†</td>
</tr>
<tr>
<td>Loos et al. (2002)</td>
<td>Belgium, F</td>
<td>238, twin pairs</td>
<td>18–34</td>
<td>SKF, WHR</td>
<td>ANOVA, R</td>
<td>+</td>
<td>–†</td>
<td>–†</td>
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<tr>
<td><strong>Middle or old age</strong></td>
<td></td>
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<tr>
<td>Gunnasdottir et al. (2004)</td>
<td>Iceland, M + F</td>
<td>3707</td>
<td>50</td>
<td>SKF</td>
<td>R</td>
<td>–F</td>
<td>–F</td>
<td></td>
</tr>
<tr>
<td>Law et al. (1992)</td>
<td>UK, M</td>
<td>1084</td>
<td>51 or 60</td>
<td>WHR</td>
<td>R</td>
<td>–†</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Sayer et al. (2004)</td>
<td>UK, M</td>
<td>737</td>
<td>64</td>
<td>BMI, SKF, WHR</td>
<td>R</td>
<td>+</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Kensara et al. (2005)</td>
<td>UK, M</td>
<td>32</td>
<td>64–72</td>
<td>DXA, BP</td>
<td>ANOVA</td>
<td>–†</td>
<td>–†</td>
<td></td>
</tr>
<tr>
<td>Gale et al. (2001)</td>
<td>UK, M + F</td>
<td>143</td>
<td>70–75</td>
<td>DXA</td>
<td>R</td>
<td>+</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

M, male; F, female; AA, arm anthropometry; BP, Bodpod; DXA, dual-energy X-ray absorptiometry; SKF, skinfold thickness; BIA, bioelectrical impedance analysis; TA, thigh anthropometry; W/H, weight-for-height; WC, waist circumference; WHR, waist:hip ratio; UWW, underwater weighing; CC, case–control study; PC, partial correlation; R, regression; TT, t test; 0, no association; +, positive association; –, negative association.

*Anthropometry not fully described.
†Results significant only after adjustment for current weight or BMI.

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unclear whether such a statistical approach is genuinely demonstrating the programming of adiposity, or simply reflects confounding as a result of the association of birth weight with lean mass.

A similar problem concerns investigations of central adiposity. Several studies using simple anthropometry have associated low birth weight with increased waist:hip ratio or subscapular:triceps skinfold thickness, but with the association often emerging only after adjustment for current weight or BMI; while several others have found no such association. The Avon Longitudinal Study of Parents and Children (Rogers et al. 2006) has demonstrated a direct rather than inverse association between birth weight and subsequent waist circumference, and no association between birth weight with truncal fat mass, measured by dual-energy X-ray absorptiometry, after adjustment for total body fat. As mentioned earlier, when associations between birth weight and body composition are dependent on adjustment for current BMI, the appropriate interpretation is that growth patterns between birth and follow up are responsible rather than birth weight itself (Lucas et al. 1999). Thus, it is plausible that these data are best interpreted as demonstrating the programming of reduced lean mass by low birth weight, such that for any given follow-up weight those individuals maintaining reduced lean mass have a higher proportion of fat in weight. This scenario, requiring statistical adjustment for current size, does not actually link low birth weight directly with increased central adiposity. Resolution of this controversy is likely to require both improved body-composition methodologies and careful statistical analyses.

The studies described earlier have all investigated associations between birth weight and body composition in childhood, and surprisingly little attention has been directed to body composition at birth itself. A particularly intriguing scenario has been reported for low-birth-weight babies from India (Yajnik et al. 2003). In comparison with normal-birth-weight babies from the UK, Indian neonates from Pune were found to have severe deficits in weight and abdominal circumference, but relatively modest deficits in skinfold thickness, especially on the trunk. The authors of the study have suggested that the Indian neonates are characterised by a ‘thin–fat’ phenotype, having reduced lean mass but preserved truncal fat, and hence a disproportionately high fat mass. At later ages Indians likewise appear to have a disproportionately high fat content for any given BMI value (Wang et al. 1994), hence there is already strong evidence of ethnic differences in physique and body composition, and the Pune data suggest such ethnic differences may derive in part from variability in fetal growth. It remains unclear whether such a thin–fat baby represents a ‘thrifty gene’ effect, possibly characteristic of Asians, or a ‘thrifty phenotype’ effect that is characteristic of low-birth-weight infants.

More broadly, associations between birth weight and later phenotype are notoriously difficult to interpret, because of the inverse association between birth weight and subsequent rate of infant weight gain. This inverse association reflects both regression to the mean (a statistical phenomenon) and catch-up (a biological phenomenon) growth. Distinguishing between regression to the mean and catch-up growth is problematic; however, recently Cameron et al. (2005) have suggested that this issue can be addressed if data are expressed in z-score format, allowing adjustment for population regression to the mean.

**Postnatal growth and subsequent body composition**

Re-evaluation of the evidence concerning the programming of adult diseases in Western populations strongly suggests that postnatal weight gain is an important component of the process (Lucas et al. 1999). For many aspects of the metabolic syndrome, risk is greatest in those born small who subsequently gain the most weight (Bavdekar et al. 1999; Adair & Cole, 2002; Barker et al. 2002). Recent studies have begun to distinguish the contributions of different periods of postnatal weight gain to later phenotype and disease risk (Barker et al. 2005), but the generalisability of such findings remains unclear.

As with birth weight, the majority of studies on infant growth rate have used BMI or skinfold thickness as the outcome. Studies are generally consistent in demonstrating a positive association between infant weight gain and later BMI or skinfold thickness (Stettler et al. 2002a,b; 2003; Ong et al. 2000). For example, a recent systematic review (Baird et al. 2005) has found that both large and rapidly-growing infants have increased risk of subsequent obesity, categorised by BMI. It is less clear, however, the extent to which these studies demonstrate a clear link between early growth and later fatness, as opposed to later size. Very few studies have adjusted childhood body fat for differences in height, even though fast infant growth is associated with greater height subsequently (Sachdev et al. 2005; Wells et al. 2005).

In the large Avon Longitudinal Study of Parents and Children cohort faster infant growth was shown to be associated with greater central adiposity as indicated by skinfold thickness (Ong et al. 2000). In the same study rapid infant growth was found to be associated with indications of growth faltering in utero, being more likely in the offspring of mothers who smoked or who had other clinical signs of fetal growth retardation. This study therefore confirms that associations between birth weight and later fat distribution are generated at least in part by the rate of postnatal growth, but are also associated with prenatal growth patterns.

In contrast to the evidence for birth weight, the effect of infant weight gain on later body composition appears to differ systematically between industrialised and developing countries (Table 2). In European studies greater infant weight gain predicts height, weight, lean mass, fat mass and waist circumference in late adolescence (Euser et al. 2005; Ekelund et al. 2006), while weight at 1 year of age likewise predicts weight, lean mass and fat mass in adults (Sayer et al. 2004). Unpublished data for UK children (approximately 200; S Chomtho, JE Williams, JCK Wells and MSFewtrell, unpublished results) link infant weight gain with later adiposity, but not with lean mass. In contrast, in three studies from non-Western populations (Li et al. 2003; Sachdev et al. 2005; Wells et al. 2005) infant
weight or BMI gain was found to be associated with later weight, height and lean mass, but not with later fat mass.

Whilst further studies are required to confirm this disparity, current evidence suggests that whether infant weight gain is directed to lean mass or fat mass is mediated by the disparity between size at birth and genetic potential. Individuals from industrialised populations, typically near their genetic potential of birth, may be unable to translate increased energy intake into greater lean mass and may be obliged to store the excess energy as fat. On the other hand, individuals from developing or modernising countries, on average small at birth, may have greater capacity to direct any additional energy directly to reducing deficits in lean mass. Consistent with this scenario, it has been shown (Ezzahir et al., 2005; Ibáñez et al., 2006) that catch-up growth in small-for-gestational-age infants is associated only with later fatness and the risk of being overweight if the catch-up persists beyond the first year of life.

Such a hypothesis might appear inconsistent with the findings from the Avon Longitudinal Study of Parents and Children cohort, for which catch-up growth was found to be associated with both indices of growth retardation and central adiposity at 5 years. However, in this study rapid growth was quantified over the first 2 years of life, whereas the studies mentioned earlier suggest that the critical window, during which weight gain following low birth weight programmes lean mass, closes within the first year of life. Thus, the findings of the Avon Longitudinal Study of Parents and Children may simply reflect the persistence of fast growth beyond this window, resulting in associations between infant weight gain and later fatness. Further studies with improved study designs and methodologies are required to investigate this issue.

More recently, several studies have focused specifically on the immediate postnatal period. In formula-fed infants weight gain in the first 8 d of life was shown to be associated with the risk of being categorised as obese in early adulthood (Stettler et al., 2005). Consistent with these data, weight gain in the first 2 weeks of postnatal life in preterm infants was found to be associated with insulin resistance during adolescence (Singhal et al., 2003a). In this randomised controlled trial those participants given a lower-nutrient diet were found to have reduced insulin resistance in adolescence, a consequence attributed to slower early growth in this group. The authors subsequently have generalised from this study to propose that ‘growth acceleration’ after birth, and particularly in the first weeks post partum, itself comprises the primary mechanism by which adult diseases, including obesity, are programmed (Singhal & Lucas, 2004).

Since the majority of centile crossing in infancy typically occurs very early on, growth acceleration could indeed link low birth weight with later disease risk. However, an alternative explanation for these data could be that the introduction of a high-nutrient diet in the immediate postnatal period programmes hormonal axes rather than growth rate per se, and that the effect on adolescent insulin resistance is derived from such hormonal programming rather than the process of growth itself. The mechanism by which growth contributes to the programming of disease therefore remains unclear, especially since initial catch-up growth seems more favourable in the populations of developing countries, and insulin resistance only appears after infancy (Ibáñez et al., 2006).

The impact of postnatal growth on later body composition is not restricted to the infant period. Weight gain during childhood also appears to exert effects on later body composition. Initial studies suggested that ‘adiposity rebound’, the age at which childhood BMI reaches a natural nadir before increasing again, was a significant predictor of later obesity (Rolland-Cachera et al., 1987). More recent studies have suggested that such rebound is both a misnomer, in that it refers to BMI rather than adiposity per se (Wells, 2000), and also a statistical artefact since the time of rebound is inherently a function of initial BMI (Cole, 2004). A more appropriate interpretation of these data is simply that those gaining weight rapidly in childhood have a significantly increased risk of obesity subsequently. It is not clear whether childhood weight gain represents programming per se, given that linear growth is by this time canalised, and any effects on fatness are theoretically reversible. Nevertheless, there are some indications that rapid rise through BMI centiles during childhood is a key contributing factor to the risk of heart disease (Barker et al., 2005).

Overall, therefore, growth patterns in both infancy and childhood contribute to the ontogenetic development of body composition and the risk of obesity. Further studies are urgently required in order to elucidate in greater detail relationships between linear growth \(v\) weight gain on the one hand, and later tissue masses and tissue distributions on the other.

### Table 2. Associations of infant weight gain with body composition indices during adolescence and young adulthood

<table>
<thead>
<tr>
<th>Reference</th>
<th>Population</th>
<th>(n)</th>
<th>Age (years)</th>
<th>Outcomes</th>
<th>Design</th>
<th>Lean</th>
<th>Fat</th>
<th>Central fat</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wells et al. (2005)</td>
<td>Brazil, M</td>
<td>172</td>
<td>9–10</td>
<td>BIA</td>
<td>ANOVA, R</td>
<td>+</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Ekelund et al. (2006)</td>
<td>Sweden, M+F</td>
<td>248</td>
<td>17</td>
<td>BP</td>
<td>R</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Euser et al. (2005)</td>
<td>The Netherlands, M+F</td>
<td>403</td>
<td>19</td>
<td>BMI, SKF, WHR</td>
<td>R</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Li et al. (2003)</td>
<td>Guatemala, M+F</td>
<td>267</td>
<td>21–22</td>
<td>SKF</td>
<td>R</td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sachdev et al. (2005)</td>
<td>India, M+F</td>
<td>1526</td>
<td>26–32</td>
<td>BMI, SKF, WHR</td>
<td>R, PC</td>
<td>+</td>
<td>+</td>
<td></td>
</tr>
</tbody>
</table>

M, male; F, female; BIA, bioelectrical impedance analysis; BP, Bodpod; SKF, skinfold thickness; WHR, waist:hip ratio; PC, partial correlation; R, regression; 0, no association; +, positive association.
Influence of nutrition

It is not possible to offer here a comprehensive account of the role of early-life nutrition in the programming of body composition. Instead, two important aspects of nutrition will be considered in order to highlight contrasting important issues.

First, there is clear evidence that fetal nutrition underlies early variability in growth but with disparity between the influence of maternal v. placental nutrient concentrations show modest or undetectable association with birth weight (Godfrey et al. 1996; Ceesay et al. 1997; Mathews et al. 1999, 2004), maternal metabolism appears to exert markedly stronger effects. The strongest evidence derives from studies of gestational maternal diabetes, a condition that develops in the second half of pregnancy and causes excessive transfer of glucose from mother to fetus. This process in turn induces hyperglycaemia and altered pancreatic structure and function in the offspring, leading to macrosomia and high body fat levels. Follow-up studies suggest an increased risk of childhood obesity (for review, see Wells, 2007). Maternal type 2 diabetes induces similar effects as discussed earlier. Similar findings emerge from studies of maternal blood pressure, with greater blood pressure associated with greater birth weight across the normotensive range (Wells, 2007).

Collectively, such research shows that aspects of increased energy transfer during pregnancy are associated with greater birth weight, and in turn with later body composition. The composition of fetal fuel supply appears key to this association, and it is plausible that stronger effects of maternal pregnancy diet may be uncovered if analyses are extended to maternal glycaemic load and/or response, rather than diet composition itself (for example, see Moses et al. 2006).

In contrast, it has proved to be more difficult to demonstrate a clear impact of early postnatal diet on later body composition. A number of studies have reported that breast-feeding is associated with a decreased risk of later obesity categorised by BMI. Two recent meta-analyses (Arenz et al. 2004; Owen et al. 2005) have supported this hypothesis, and have suggested both categorical and dose–response protective effects. However, studies that have focused on body composition itself as the outcome have in general failed to support this hypothesis. A large study of adult males from Brazil (Victora et al. 2003) has found no association between breast-feeding and adult body fat in men, while an analysis from the Avon Longitudinal Study of Parents and Children cohort (Toschke et al. 2007) has found a negligible effect of breast-feeding on mean body fat. Although this study has found a small protective effect against being in the top decile of body fat, this effect was shown to be greatly attenuated after adjusting for confounders, reducing any physiological effect attributable to diet itself.

The complex scenario relating to breast-feeding highlights several major difficulties when investigating the role of nutrition in the programming of body composition. First, detailed information about breast-feeding practice is rarely available in large cohort studies. Second, reverse causation may contribute, with those experiencing difficulties breast-feeding resorting to ‘aggressive’ formula feeding in order to overcome early deficits in the weight gain of their infants. Third, breast-feeding is associated with a variety of social factors, and it is difficult to be confident that diet, as opposed to associated factors, explains subsequent outcome. More generally, randomised controls of early diet are rare, particularly in relation to fetal nutrition, and in the case of breast-feeding are unethical.

Identifying causal roles of nutrition in the programming process is therefore difficult. Nevertheless, the importance of this issue should not be denied, since early-life diet remains implicated in the aetiology of obesity and public health policy requires reliable evidence of this role in order to recommend appropriate dietary practices. Important components of early-life nutrition include not only maternal metabolism and infant feeding mode but also postnatal feeding for those infants born preterm, feeding schedule in the hours following birth and weaning schedule. As with maternal nutrition during pregnancy, infant dietary glycaemic load may well prove to be of importance.

However, it is noted that despite the considerable emphasis directed to the notion of fetal ‘undernutrition’ as a key factor in the programming of disease, maternal diet during pregnancy appears to exert relatively little effect on offspring birth weight. This small effect is in marked contrast to the impact of maternal pregnancy metabolism (Wells, 2007). Given the rapidly-increasing prevalence of the metabolic syndrome in many populations, reflected in increasing prevalences of obesity, diabetes and hypertension in pregnant mothers, the influence of maternal pregnancy metabolism, as opposed to dietary intake, on offspring body composition merits further attention.

Evolutionary perspective

The fact that early growth rate predicts later risk of disease might be interpreted as deriving from interactions between human biology and the Western industrialised niche that the human population has entered only recently. For example, CVD represents a relatively new cause of mortality, rarely reported before the 20th century. It would be more appropriate, however, to consider early-life programming as an integral and fundamental component of human biology, and the role of body composition in this process clarifies this position. To focus disproportionately on the ‘thrifty phenotype’ of low-birth-weight infants as a risk factor for later disease is to miss broader biological patterns, since associations between birth weight and later outcome tend to hold broadly across the entire range of birth weight. When adopting an evolutionary perspective it is helpful to discard the terminology of ‘programming’ and to refer instead to the ‘environmental induction’ of phenotype (Bateson, 2001). Such an approach avoids incorrectly suggesting that early environmental conditions in themselves contain specific cues for disease.

All offspring should be regarded as subject to environmental phenotypic induction during early development, with growth trajectory (and later health profile) inevitably
being associated with maternal phenotype before conception and during pregnancy and lactation (Wells, 2003). The fact that the two genders show different associations between early growth and later body composition is a further indication that such phenotypic induction is a normal component of human development. It is the interaction between such normal developmental plasticity and relatively-modern environmental factors that is now generating clear associations between early experience and later disease risk.

Evolution has been described as a game in which the reward for genes ‘winning’ (surviving) in one time period is the chance to ‘play again’ in the next (Slobodkin & Rappaport, 1974). Although it is commonly assumed that natural selection acts on traits (e.g. eye colour, height), it is more appropriate to consider strategies as the true target of selection (Houston & McNamara, 1999). Natural selection has clearly favoured substantially higher levels of body fat in adult females, and of lean mass in adult males (Fig. 2). Such differences have been attributed to the differing energy requirements and behaviours associated with reproductive fitness, with the greater fatness of females aiding the high energy costs of pregnancy and lactation, and the greater lean mass of males plausibly aiding intermale competition or resource acquisition (Wells, 2006).

The data reviewed earlier indicate that even in fetal life male and female offspring have begun to pursue different strategies in their relative allocation of energy to somatic and reproductive fitness, with the greater fatness of females plausibly aiding intermale competition rather than nutritionally regulated. It has been argued previously (Wells, 2003) that such canalisation has evolved to buffer the greater variability in energy supply that occurs with weaning. In more-modern environments, however, insulin resistance can interact with a high-energy-density diet throughout childhood, resulting in excess fat accumulation from the time point when growth becomes canalised and lean mass loses its plasticity.

### Priorities for future research

A major limitation of existing work on the programming of body composition during early life has been hindered by a paucity of appropriate techniques. Most body-composition techniques are either impractical for infants or of unknown accuracy. Isotope dilution has been used in a number of studies of infants for the last two decades, but follow-up studies have been rare. There are now several techniques that can be used to measure body composition in early life, which include, in addition to isotope dilution, whole-body air-displacement plethysmography and MRI scanning. These techniques should now be used to investigate: first, body composition at birth, and its association with fetal experience; second, body composition during infancy and the effects of variability in growth rate. Such work is likely to make a major contribution to the understanding of the process by which early environmental factors are associated with the risk of the metabolic syndrome, and this research may be particularly important in understanding ethnic disparities in disease risk.

### Summary

The ontogenetic development of body composition is implicated in several ways in the aetiology of the metabolic syndrome. Lean mass has implications for glucose uptake and physical capacity for work and exercise. Fat mass and its distribution has implications for reproductive function and risk of disordered metabolism. Recent studies have invoked a shift from undue emphasis on the consequences of low birth weight to a broader appreciation of the long-term effects of growth throughout the developmental period and across the entire range of body size. Associations between growth and later body composition strongly implicate nutrition as the underlying mechanism.
However, existing evidence remains sparse, and in the case of breast-feeding, inconsistent. A new focus on body composition itself in early life is likely to make a major contribution to the understanding of the programming of the metabolic syndrome.

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