Birth weight, weight at one year, and left ventricular mass in adult life

M Vijayakumar, C H D Fall, C Osmond, D J P Barker

Abstract

Objective—To examine how fetal and infant growth are related to left ventricular mass in adult life.

Design—A follow up study of men born during 1920–30 whose birth weights and weights at 1 year were recorded.

Setting—Hertfordshire, England.


Main outcome measure—Left ventricular mass calculated from measurements of interventricular septal thickness and left ventricular posterior wall thickness and left ventricular internal diameter at end diastole measured by M mode echocardiography.

Results—Left ventricular mass was highest in men with the lowest weight at 1 year and fell with increasing weight at 1 year ($r = 0.18$, $P = 0.01$). Left ventricular mass was not related to birth weight. The relation with weight at 1 year was independent of factors in adult life known to influence left ventricular mass, including body size, systolic blood pressure, and age. The enlarged left ventricular mass associated with reduced growth in infancy was concentric, affecting both the interventricular septum and the left ventricular posterior wall. Concentric left ventricular hypertrophy is known to be associated with increased death rates from coronary artery disease.

Conclusion—Low weight at 1 year is associated with concentric enlargement of the left ventricle in adult life. This is consistent with a previous finding of higher mortality from cardiovascular disease in men of low weight at 1 year, and provides further evidence that cardiovascular disease may be partly programmed in early life. The left ventricular enlargement may be a long term result of haemodynamic changes in utero or of persisting changes in growth factor concentrations.

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Keywords: birth weight; left ventricular enlargement; weight at one year; cardiovascular disease

Recent findings suggest that the pathogenesis of coronary heart disease begins in fetal life and infancy. Among 10 141 men born during 1911–30 in Hertfordshire, England, whose birth weights and weights at 1 year were recorded, men with the lowest birth weights and weights at 1 year had the highest death rates from coronary heart disease. Reduced growth in utero and during infancy was also associated with an increased risk of hypertension, non-insulin dependent diabetes, and raised LDL cholesterol and fibrinogen concentrations in adult life. These findings have led to the hypothesis that coronary heart disease originates from early programming whereby undernutrition during sensitive periods in early development permanently changes the body's structure and physiology.

Left ventricular hypertrophy, determined by either electrocardiography or echocardiography, has been shown to be a strong predictor of morbidity and death from coronary heart disease. Much of the variance in left ventricular mass in adults is unexplained by known influences, such as body size and systolic blood pressure. The human heart has its highest growth rates in fetal and early postnatal life. Neither the relation between this early growth and adult left ventricular mass nor the long-term effects of early growth retardation have yet been studied. Doppler ultrasound studies have shown that in some growth retarded fetuses haemodynamic changes result in the redistribution of blood flow to the brain at the expense of the viscera and lower limbs. This redistribution is associated with an increase in left ventricular blood flow and increased peripheral arterial resistance. Such haemodynamic changes could produce long-term changes in left ventricular size.

We measured left ventricular mass in a sample of men whose birth weights and weights at one year had been recorded.

Patients and methods

In Hertfordshire from 1911 onwards each birth was notified by the attending midwife, and a health visitor saw the child periodically throughout infancy. The child's birth weight and weight at one year were recorded. We used these records to trace men who were born in the county between 1911 and 1930, and to determine cardiovascular mortality rates. We subsequently examined a sub-sample of the men who were born in East Hertfordshire between 1920 and 1930, and still live there. Their occupation and their father's occupation at the time of their birth were used to determine social class, currently and at birth. Their blood pressure, serum lipids, plasma clotting factors, and glucose
and insulin concentrations were measured, and the results have previously been reported. We reapproached the 370 men who had complete measurements on all blood samples and asked them to take part in this study. Seven of the 370 had died and 11 had moved away: 290 (82%) of the remaining 352 took part. The men were visited at home by a nurse who asked for details of current smoking habits and alcohol consumption. Alcohol consumption was converted to the total number of units per week (1 unit = 10 ml ethanol).

After the interview subjects were asked to come to a local clinic, where left ventricular mass was measured echocardiographically according to the recommendations of the American Society of Echocardiography. Cross-sectionally directed M mode echocardiographic examination was carried out by one of two doctors (MV and CHDF), with the subjects in the partial left decubitus position, using a 2-25 Hz transducer (Siemens 5000, Esota Biomedica, Italy). The interventricular septum and left ventricular posterior wall, at or just below the mitral valve leaflets, were simultaneously visualised throughout the cardiac cycle. The M mode cursor was placed perpendicularly to the long axis of the left ventricle. M mode prints were made using a video printer (Sony UP930, Japan). Measurements of the left ventricular internal diameter at end diastole (LVIDD) and end systole (LVIDS), interventricular septal thickness at end diastole (IVSD), and left ventricular posterior wall thickness at end diastole (LVPW) were made from the prints using a digitiser (Genius GT1212B, Taiwan). Measurements were made in five cardiac cycles and the mean was used in the analysis.

Left ventricular mass was calculated using the formula:

\[ LV \text{ mass} = 0.8 \times (IVSD + LVIDD + LVPW) - LVIDS + 0.6 \times \text{height} \]

Relative wall thickness (RWT) was calculated using the formula:

\[ \text{RWT} = \frac{2 \times (LVPW/LVIDD)}{\text{RWT}} \]

Left ventricular volume was calculated using Teichholz formula, and the ratio between left ventricular mass and left ventricular volume at end diastole (M/V ratio) was derived. The RWT and M/V ratio are indicators of the concentricity of left ventricular hypertrophy. Cardiac output (CO) was calculated as (stroke volume \times heart rate).

The men's weight was measured on a Seca scale and their height was measured on a portable stadiometer (CMS Instruments Ltd). Body surface area (m²) was calculated using the formula:

\[ 0.007184 \times \text{weight (kg)}^{0.425} \times \text{height (cm)}^{0.725} \]

Systolic and diastolic blood pressures were measured with an automated device (Dinamap, Critikon, Ascot) and a cuff of the appropriate size. The Rose/WHO chest pain questionnaire was administered. A 12 lead electrocardiograph was obtained according to the 1982 Minnesota protocol. Electrocardiographs were coded by two trained coders who were blind to the infant data. Men were defined as having coronary heart disease if one or more of the following were present: a history of typical angina according to the Rose/WHO chest pain questionnaire, ECG Minnesota codes 1–1 or 1–2 (major Q waves), or a history of coronary artery revascularisation surgery.

**STATISTICAL METHODS**

The distributions of left ventricular mass and M/V ratio were skewed. We transformed the values using logarithms. Differences in left ventricular mass in men with and without coronary heart disease were assessed by a two sample t test, and differences between smokers, non-smokers, and ex-smokers were assessed by one way analysis of variance. Multiple linear regression analysis was used to analyse the relation of left ventricular mass with birth weight, weight at one year, factors in adult life known to influence ventricular mass including body size, age, and systolic blood pressure and other potential confounding factors including social class (treated as an ordinal variable) and alcohol consumption.

**Results**

Of the 290 men examined we excluded five with valvar heart disease and two with left ventricular dyskinesia. In a further 81 (29%) men suitable views of the left ventricle were not obtained: in 32 this was because an inadequate parasternal echocenic window made it impossible to align the M mode cursor perpendicularly to the long axis of the left ventricle and in 49 either the posterior septal wall or the posterior ventricular wall was not seen distinctly enough to make measurements. The analysis is based on 202 men. Their mean age was 66.9 (SD 3.2) years.

Their geometric mean left ventricular mass was 203 g (range 103–377 g; geometric standard deviation 1.26). Left ventricular mass was higher in men of larger body size, rising with increasing body mass index (weight (kg)/height(m)²) (r = 0.25; P = 0.0003), and with increasing body surface area (r = 0.21; P = 0.002). It also rose with increasing systolic blood pressure (r = 0.27; P = 0.0001) and with increasing age (r = 0.13; P = 0.07). Geometric mean ventricular mass was higher (r = 0.26 g) in the 33 men with coronary heart disease than in the men without (199 g) (difference 27 g; 95% C.I. 8 to 47; P = 0.004). There were no significant differences in left ventricular mass between smokers (geometric mean 203 g, n = 30), ex-smokers (geometric mean 205 g, n = 136), and lifetime non-smokers (geometric mean 198 g, n = 36). Left ventricular mass was not related to alcohol consumption, current social class, or social class at birth.

Table 1 shows mean left ventricular mass according to the men's weight at one year (see also figure) and birth weight. Values in table 1 are shown both unadjusted and adjusted for surface area, the measure of body size against which left ventricular mass is traditionally indexed. Left ventricular mass was highest among men in the lowest group of weight at one year, and fell progressively with increasing weight at one year (r = 0.18, P = 0.01). Left
Table 1  Geometric mean left ventricular mass according to weight at age one year and birth weight

<table>
<thead>
<tr>
<th>Age (y) at one year</th>
<th>Mean left ventricular mass (g)</th>
<th>Mean left ventricular mass (g) adjusted for BSA</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>418</td>
<td>418</td>
</tr>
<tr>
<td>2</td>
<td>423</td>
<td>423</td>
</tr>
<tr>
<td>3</td>
<td>425</td>
<td>425</td>
</tr>
<tr>
<td>4</td>
<td>426</td>
<td>426</td>
</tr>
</tbody>
</table>

Birth weight (lb):

<table>
<thead>
<tr>
<th>Weight at one year</th>
<th>Mean body surface area (m²)</th>
<th>Mean body surface area (m²) adjusted for BSA</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤2</td>
<td>198 (25)</td>
<td>198 (25)</td>
</tr>
<tr>
<td>&gt;2</td>
<td>172 (16)</td>
<td>172 (16)</td>
</tr>
</tbody>
</table>

All figures in parentheses are numbers of men.

Table 3  Multiple regression analysis of log left ventricular mass (g) with body surface area, systolic blood pressure, age, and weight at one year

<table>
<thead>
<tr>
<th>Variable</th>
<th>Regression coefficient</th>
<th>SE</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Body surface area (m²)</td>
<td>0.0450</td>
<td>0.0170</td>
<td>0.0002</td>
</tr>
<tr>
<td>Systolic blood pressure (mm Hg)</td>
<td>0.0027</td>
<td>0.0008</td>
<td>0.001</td>
</tr>
<tr>
<td>Age (y)</td>
<td>0.0047</td>
<td>0.0048</td>
<td>0.3</td>
</tr>
<tr>
<td>Weight at one year (lb)</td>
<td>-0.2005</td>
<td>0.0004</td>
<td>0.002</td>
</tr>
<tr>
<td>Constant</td>
<td>4.2094</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

y variable = log left ventricular mass (g).

Table 4  Echocardiographic measurements of the left ventricle and calculated values of relative wall thickness, M/V ratio and cardiac output according to weight at one year

<table>
<thead>
<tr>
<th>Variable</th>
<th>Weight at one year (pounds)</th>
<th>P value* for trend allowing for current BSA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measured values:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Left ventricular internal diameter at end diastole (LVIDD) (mm)</td>
<td>46-1 45-3 45-7 46-3 45-6 47-4 46-0 51-1 0.4 0.9</td>
<td></td>
</tr>
<tr>
<td>Interventricular septal thickness at end diastole (IVSTD) (mm)</td>
<td>24.2 23.9 24.8 25.3 23.9 25.1 24.7 4.3 0.6</td>
<td>1.0</td>
</tr>
<tr>
<td>Left ventricular posterior wall thickness at end diastole (LVPWD) (mm)</td>
<td>15-1 14-2 14-3 13-6 13-5 12-8 13-9 2.5 0.009</td>
<td>0.003</td>
</tr>
<tr>
<td>Calculated values:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Relative wall thickness (RWT)</td>
<td>0.53 0.47 0.48 0.44 0.44 0.42 0.46 0.1 0.004</td>
<td>0.007</td>
</tr>
<tr>
<td>M/V ratio (g/ml)</td>
<td>2.5 2.2 2.2 2.0 2.0 1.9</td>
<td>2.1</td>
</tr>
<tr>
<td>Cardiac output (CO) (l/min)</td>
<td>4.3 4.6 4.8 4.9 4.7 5.0 4.8 1.3 0.2</td>
<td>0.6</td>
</tr>
</tbody>
</table>

*Geometric standard deviation. †Multiple linear regression.
fibroinogen and serum LDL cholesterol concentrations.

Discussion
We have found that low weight at one year was associated with increased left ventricular mass in adult life (table 1). The relation was statistically significant, and independent of current body size (table 2) and systolic blood pressure (table 3). The increased left ventricular mass reflected concentric enlargement, affecting both the interventricular septum and the left ventricular posterior wall. Concentric left ventricular enlargement is a known predictor of coronary heart disease morbidity and mortality. The Framingham study showed that in men aged 40 years and over, a 50% increase in left ventricular mass was associated with a doubling in incidence of cardiovascular disease. Left ventricular mass was not related to birth weight. We have previously shown that in men weight at one year is a stronger predictor of mortality from coronary heart disease than birth weight. Our study was confined to men born and still living in East Hertfordshire, and willing to take part. This introduced the possibility of selection bias. However, our analysis was based on comparisons within the sample, and bias would be introduced only if the relation between early growth and coronary heart disease were different in those studied and not studied. This seems unlikely. It was not possible to obtain adequate echocardiographic views for measurement of left ventricular mass in 29% of the men. This is a recognised problem, especially in the elderly, and our success rate was similar to other studies in this age group. There were no differences in age, body mass index, body surface area, systolic blood pressure, birth weight or weight at one year between men for whom we did or did not measure left ventricular mass.

Although growth during infancy may be influenced by feeding, the relation of coronary heart disease mortality to weight at one year is independent of method of feeding in infancy. Undernutrition in late pregnancy may be followed by growth failure in infancy. At birth the baby may be short but have a normal birth weight. Raised plasma fibrinogen concentrations in adult life are associated with shortness at birth, and low weight at one year, but not with low birth weight. A common origin in undernutrition in late gestation could explain the association between left ventricular mass and raised plasma fibrinogen in our study. A similar explanation could account for the association between left ventricular mass and LDL cholesterol.

One link between undernutrition in late gestation and persisting increase in left ventricular mass could be mediated through cardiovascular adaptive changes in utero. In late gestation the human fetus may respond to nutrient deprivation by maintaining the brain growth at the expense of the growth of the trunk. The redistribution of blood flow that accompanies this is associated with an increase in peripheral resistance, a reduction in blood flow in the abdominal aorta, and an increase in left ventricular blood flow. These changes may lead to permanent left ventricular enlargement.

Another possible link between undernutrition in late gestation and left ventricular mass is persisting changes in hormones which regulate growth. We have previously suggested that failure of infant growth is associated with defects in the growth hormone/insulin-like growth factor axis. These may influence cardiovascular structure. Both growth hormone and insulin-like growth factor-1 stimulate ventricular growth, effects which are evident in acromegaly.

We thank the men who took part in the study, the staff of the Hertfordshire Family Health Services Authority who helped to trace the men. The fieldwork was coordinated by Mrs P Harwood. Nurses S Haynes, P Howell, R Rosenthal, and S Wolfe carried out the home interviews and helped with the clinics. ECGs were coded by Mrs C Rose and Mrs N Keen. M V was a Commonwealth Research Fellow. The study was funded by the Medical Research Council.

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