Associations between birth weight, early childhood weight gain and adult lung function

R J Hancox,1 R Poulton,1 J M Greene,2 C R McLachlan,1 M S Pearce,3 M R Sears2

ABSTRACT

Background: Low birth weight is associated with lower values for spirometry in adults but it is not known if birth weight influences other measures of pulmonary function. It is also unclear whether postnatal growth affects adult lung function. The associations between birth weight, postnatal growth and adult lung function were assessed in an unsellected birth cohort of 1037 children.

Methods: Birth weight, weight gain between birth and age 3 years, and lung function at age 32 years were measured. Analyses were adjusted for adult height and sex and further adjusted for multiple other potential confounding factors.

Results: Birth weight was positively correlated with spirometric (forced expiratory volume in 1 s and forced vital capacity) and plethysmographic (total lung capacity and functional residual capacity) lung function and with lung diffusing capacity. These associations persisted after adjustment for confounding factors including adult weight, exposure to cigarette smoke in utero and during childhood, personal smoking, socioeconomic status, asthma and gestational age. Weight gain between birth and age 3 years was also positively associated with lung diffusing capacity, and with higher values of lung volumes in men after adjustment for covariates. Neither birth weight nor postnatal weight gain was associated with airflow obstruction.

Conclusions: Low birth weight and lower weight gain in early childhood are associated with modest reductions in adult lung function across a broad range of measures of lung volumes and with lower diffusing capacity. These findings are independent of a number of potential confounding factors and support the hypothesis that fetal and infant growth is a determinant of adult lung function.

The “fetal origins” hypothesis proposes that impairment of intrauterine growth may have long term consequences for physiological function and risk for adult disease.1 With respect to respiratory health, a number of studies have reported an association between low birth weight and lower lung function in adulthood.2–6 Although not all studies have observed this association,7–8 a recent meta-analysis and subsequent reports support a modest positive association between birth weight and adult forced expiratory volume in 1 s (FEV1).9–11 These findings support the hypothesis that intrauterine growth is a critical time for lung development and a determinant of eventual lung function.

Whether fetal growth influences other aspects of respiratory function is not known because most studies have relied on spirometry to assess adult lung function. Studies of other measures of lung function have been small and restricted to patients with severe prematurity who have developed bronchopulmonary dysplasia.12 Even for spirometric outcomes, it remains uncertain whether low birth weight is primarily associated with an obstructive or restrictive pattern of reduced lung function in adulthood. Although several studies have found parallel reductions in both FEV1 and forced vital capacity (FVC) indicating restriction,9 10 11 13 other studies have not found a significant effect on FVC, suggesting an obstructive defect.2 One potential reason for the inconsistency between studies is that some have been retrospective while others have been unable to control for factors that may confound or mediate the association between birth weight and adult lung function. For example, lower socioeconomic status and maternal smoking may cause both low birth weight and impaired postnatal lung growth.14 15

There is less information on whether postnatal growth influences adult lung function. One study found that adult FEV1 and FVC were positively associated with weight gain in the first year of life although the effect on FEV1 was not statistically significant in men.16 However, another cohort found that growth in the first year of life was not related to adult lung function.17 Other studies have reported that greater postnatal weight gain is associated with lower lung function in infancy.17 18

To our knowledge, there have been no studies of the association between fetal development and a comprehensive assessment of adult lung function in a population based sample. There is good reason to postulate that the impact of fetal growth on respiratory physiology may vary because the pattern of growth differs for different lung structures: whereas the number of conducting airways is normally complete at birth, 85% of alveoli are believed to develop postnatally.19 We explored the association between birth weight, postnatal growth and adult lung function in a prospective population based cohort followed to age 32 years.

METHODS

Study members were born in Dunedin, New Zealand, between April 1972 and March 1973.20 21 A total of 1087 children (91% of eligible births; 52% male) participated in the first follow-up assessment at age 5 years, constituting the base sample for the remainder of the study. This analysis explores the association between birth weight, weight change between birth and age 5 years, and lung function at age 32 years, which was measured in 94% of living study members (953/1015). Study members were mostly of New Zealand/European ethnicity.

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At age 32 years, 6% identified themselves as Maori. Few study members identified with other ethnicities. The Otago Ethics Committee approved the study. Written informed consent was obtained.

Birth weight was measured for all study members. Gestational age was calculated from the date of the last menstrual period when this was recalled with confidence (n = 915). Ninety-seven per cent of children were reassessed within 1 month of their third birthday, but 31 were assessed during the second month after their birthday. Weight at age 3 years was available for 951 study members. Childhood socioeconomic status was calculated as the average of the higher socioeconomic status level of either parent, based on the education and income associated with their occupation, assessed repeatedly from the study member’s birth through to age 15 years.22 23

Information obtained about respiratory health throughout the life course was updated at age 32 years, including questions from the American Thoracic Society and the European Community Respiratory Health Survey questionnaires.24 25 Current asthma is defined as a diagnosis of asthma with episodes of asthma, wheezing or asthma medication use in the previous year. Cumulative smoking was calculated as the number of pack years cigarettes smoked up to age 32 years (20 cigarettes per day for 1 year = 1 pack year). Smoking history for both parents/guardians was obtained from the adult attending with the study member at ages 7, 9 and 11 years and from the study members themselves at age 13 years.26 Maternal smoking during pregnancy was recorded retrospectively at the age 9 assessment and this information was available for 777 study members.

Height and weight were measured in light clothing without shoes. Spirometry (FEV1 and FVC), static lung volumes (total lung capacity (TLC), functional residual capacity (FRC), residual volume (RV)) and single breath diffusing capacity for carbon monoxide (TlCO) and alveolar volume by methane dilution (Va) were measured to American Thoracic Society standards27–29 using a Sensormedics 6200 plethysmograph with Vmax version 4.3b software (Yorba Linda, California, USA). A portable spirometer (Spiropro, Sensormedics, Yorba Linda) was used to test those who refused to sit in the plethysmograph or were unable to attend the research unit (n = 27). All tests were reviewed by a senior technician to ensure only acceptable and reproducible results were entered for analysis. Equipment was calibrated daily, and weekly quality control measures were obtained to ensure accuracy and precision of the test equipment.

Haemoglobin was measured on a Sysmex XE2100 automated haematology analyser (Sysmex Corporation, Japan). Exhaled carbon monoxide was measured twice using a Micro CO monitor, (Micromedical, UK) and the average was recorded.

Associations between birth weight and adult lung function measurements were analysed by multiple linear regression using lung function as the dependent variable and birth weight as the main predictor (independent variable). In addition to the lung function variables listed above, FEV1/FVC and TlCO/Va ratios were analysed as indicators of airflow obstruction and the diffusing capacity per unit alveolar volume, respectively. Initial analyses of the association between birth weight and adult lung function adjusted for adult height and sex. Analyses of TlCO and TlCO/Va additionally adjusted for haemoglobin and exhaled carbon monoxide. Analyses also tested sex*birth weight interaction terms to assess if the association between birth weight and adult lung function differed for males and females. These analyses were repeated using weight change between birth and age 3 years as the main predictor.

Further analyses included both birth weight and weight change between birth and age 3 years as the main predictors. These analyses also adjusted for a number of potential covariates: gestational age, maternal smoking during pregnancy, parental smoking during childhood, personal smoking up to age 32 years in pack years, childhood socioeconomic status, current asthma at age 32 years and body weight at age 32 years. Interaction terms between birth weight and weight change up to age 3 years were calculated to assess whether the influence of either predictor depended on the other (eg, low birth associated with greater postnatal weight gain).

Women who were pregnant at the time of the age 32 assessment (n = 31) were excluded from all analyses. Visual inspection of the residuals from the regression models identified one clear outlier. This individual was excluded from the analyses. Analyses were performed using Stata 10.0 (StataCorp, College Station Texas, USA).

### RESULTS

The birth characteristics of the study members are shown in table 1. Birth weight correlated with gestational age (r = 0.46, p<0.0001) and tended to be higher in boys (mean (SD) 3.43 (0.53) vs 3.31 (0.51) kg in girls; p<0.001). Birth weight also correlated with weight at age 32 years (sex adjusted correlation: r = 0.17, p<0.001). Weight gain between birth and age 5 years was greater in boys than girls (means (SD) 11.6 (1.6) and 11.1 (1.5) kg, respectively; p<0.0001) and was weakly correlated with birth weight (sex adjusted correlation: r = 0.06, p = 0.047). Weight gain between birth and age 5 years also correlated with adult weight (sex adjusted correlation: r = 0.51, p<0.0001).

Mean values for the lung function tests are shown in table 2. Higher birth weights tended to be associated with higher spirometric and plethysmographic lung volumes in adulthood (table 3). These trends were not statistically significant in the
sex and height adjusted models but were significant in the fully adjusted analyses. Birth weight was also significantly associated with diffusing capacity (TlCO). Similar trends were found for diffusion capacity adjusted for alveolar volume (TlCO/Va) although this was not statistically significant in the fully adjusted model. Sex × birth weight interaction terms were not significant with the exception of TlCO (p = 0.023). The association between birth weight and TlCO was stronger in males (coefficient (95% CI) 1.47 (0.60 to 2.34); p = 0.001) than in females (0.36 (−0.28 to 1.00); p = 0.27) after adjustment for height, haemoglobin and exhaled carbon monoxide. This sex interaction was not apparent in the fully adjusted model when the coefficients were similar for males and females (1.19 (−0.08 to 2.46) and 1.13 (0.22 to 2.05), respectively).

Weight gain between birth and 3 years was not significantly associated with any of the spriometeric or plethysmographic lung volumes in the sex and height adjusted analyses; however, there was a trend to lower FRC values with greater weight gain (table 4). Greater weight gain was associated with higher values for adult FRC in the fully adjusted model with trends to higher (table 4). Greater weight gain was associated with higher values for TlCO and TlCO/Va in the sex and height adjusted analyses; however, coefficients represent the difference in lung function associated with a 1 kg difference in birth weight. FEV1, forced expiratory volume in 1 s; FRC, functional residual capacity; FVC, forced vital capacity; RV, residual volume; TLC, total lung capacity; TlCO, single breath diffusing capacity for carbon monoxide; Va, alveolar volume.

### Table 3 Regression analyses of age 32 lung function on birth weight

<table>
<thead>
<tr>
<th></th>
<th>Adjusted for sex and height</th>
<th>Fully adjusted</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>No</td>
<td>Coeff (95% CI)</td>
</tr>
<tr>
<td>FEV1 (ml)</td>
<td>848</td>
<td>2.0 (−22 to 26)</td>
</tr>
<tr>
<td>FVC (ml)</td>
<td>848</td>
<td>15.8 (−9 to 44)</td>
</tr>
<tr>
<td>FEV1/FVC (%)</td>
<td>848</td>
<td>−0.18 (−0.47 to 0.12)</td>
</tr>
<tr>
<td>TLC (ml)</td>
<td>816</td>
<td>16.8 (−18 to 51)</td>
</tr>
<tr>
<td>FRC (ml)</td>
<td>817</td>
<td>−30.7 (−63 to 1)</td>
</tr>
<tr>
<td>RV (ml)</td>
<td>816</td>
<td>3.4 (−14 to 21)</td>
</tr>
<tr>
<td>TlCO (ml/min/mm Hg)</td>
<td>775</td>
<td>0.22 (0.01 to 0.43)</td>
</tr>
<tr>
<td>TlCO/Va (ml/min/mm Hg/l)</td>
<td>775</td>
<td>0.01 (−0.02 to 0.05)</td>
</tr>
<tr>
<td>Va (ml)</td>
<td>825</td>
<td>22.1 (−12 to 56)</td>
</tr>
</tbody>
</table>

All analyses adjust for sex and (except for FEV1/FVC) height at age 32 years. All analyses of TlCO and TlCO/Va adjust for exhaled carbon monoxide and blood haemoglobin. Analyses in the fully adjusted models also adjust for weight gain between birth and age 3 years, gestational age, maternal smoking during pregnancy, parental smoking during childhood, personal smoking history to age 32 years, childhood socioeconomic status, current asthma at age 32 years and weight at age 32 years. Coefficients represent the difference in lung function associated with a 1 kg difference in birth weight. All analyses adjust for sex and (except for FEV1/FVC) height at age 32 years. All analyses of TlCO and TlCO/Va adjust for exhaled carbon monoxide and blood haemoglobin. Analyses in the fully adjusted models also adjust for weight gain between birth and age 3 years, gestational age, maternal smoking during pregnancy, parental smoking during childhood, personal smoking history to age 32 years, childhood socioeconomic status, current asthma at age 32 years and weight at age 32 years. Coefficients represent the difference in lung function associated with a 1 kg difference in birth weight. FEV1, forced expiratory volume in 1 s; FRC, functional residual capacity; FVC, forced vital capacity; RV, residual volume; TLC, total lung capacity; TlCO, single breath diffusing capacity for carbon monoxide; Va, alveolar volume.
DISCUSSION

In this population-based cohort, higher birth weight was associated with higher values of adult lung function. This association was not only found for spirometric lung volumes, as previously reported, but also for measures of static lung volume measured by plethysmography and for lung diffusing capacity. These findings support the hypothesis that impaired intrauterine growth may have long-term consequences for lung development.

The associations between lower birth weight and adult lung function were stronger in the fully adjusted analysis which included adjustment for adult weight. This indicates that the associations between birth weight and lung function were not mediated by the correlation between birth weight and subsequent weight gain to adulthood. However, an alternative interpretation is that it is the change between birth and adult weight which is the primary predictor—in other words, "centile crossing" as low birth weight babies catch up growth with their peers. This seems an unlikely explanation for our findings. Although birth weight was weakly correlated with weight gain between birth and age 3 years, the associations between birth weight and lung function were independent of this. There were no significant interactions between birth weight and either postnatal weight gain or adult weight for any of the lung function outcomes, indicating that the associations between birth weight and adult lung function were not contingent on subsequent growth. Finally, when the fully adjusted analyses were repeated without adjustment for adult weight, a broadly similar pattern of findings was observed (see table 3 online).

The adjusted analyses also indicate that the associations between birth weight and lung function were independent of a number of potential covariates which may have confounded or mediated the association. For example, low gestational age, lower socioeconomic status and maternal smoking have been associated with both low birth weight and impaired lung development. In fact, in our analyses, none of these was a significant independent predictor of adult lung function. Analyses excluding these covariates showed a similar pattern of findings (see table 4 online).

Interpreting the association between postnatal growth and adult lung function is more difficult. The finding that postnatal weight gain predicted adult diffusing capacity is in keeping with the fact that most alveoli are formed during this time. Weight gain between birth and age 3 years was also associated with higher values for static lung volumes in the fully adjusted analyses, which is consistent with this being the main stage of alveolar development. However, these associations were only apparent if the analyses adjusted for adult body weight (see table 4, and online table 5). This suggests that these associations may be due to confounding by the association between weight gain up to age 3 years and body weight at age 32 years which was inversely associated with all dynamic and static measures of lung volumes. Unfortunately, we do not have measurements of weight earlier in infancy to further clarify the relation between postnatal growth and adult lung volumes.

To our knowledge, this is the first report of the association between birth weight, weight gain in early childhood and a comprehensive assessment of adult respiratory function in a population-based cohort. Previous reports have been restricted to small numbers of severely premature babies at risk of bronchopulmonary dysplasia. We found that although birth weight correlated with gestational age, the associations were independent of this indicating that the effects on lung function were not due to premature birth. Excluding the small number of babies born before 37 weeks gestation made little difference to the associations between birth weight and adult lung function (data not shown).

We found little evidence for an association between either birth weight or weight gain in the first 3 years and the FEV1/FVC ratio. Although an inverse association between weight gain and the FEV1/FVC ratio was identified in men in the fully adjusted analysis, this was because weight gain was associated with higher values for FVC rather than lower values for FEV1 (data not shown). Our findings are consistent with other reports and suggest that the impairment in lung function associated with a low birth weight is compatible with a restrictive defect rather than airways obstruction. This is also true for the pattern of changes associated with lower postnatal growth in the fully adjusted analyses.

Birth weight has been used as the measure of fetal growth in many studies of the fetal origins hypothesis, including most of those analysing the association with adult lung function. Possibly this is because birth weight is the most available measure in many datasets. One cohort used multiple measures including BMI at birth and 1 year to predict adult spirometry. In keeping with this report, we found birth weight to be an equivalent or better predictor of adult lung function than some other measures of birth size, including length. Analyses using BMI at birth and change in BMI between birth and age 3 years were similar to the birth weight and weight change analyses but in some cases birth BMI appeared to be a better predictor (tables 1 and 2 online).

The study has a number of strengths. It is a large population-based cohort with a high rate of follow-up. Lung function was measured by experienced technicians with rigorous quality control measures. Information about birth weight and other perinatal factors, smoking, socioeconomic status and asthma were obtained prospectively. Even though missing data for some of the covariates reduced the numbers available for the fully adjusted analyses, the proportion of participants available for these analyses compares favourably with other studies. Moreover, there were no significant differences in birth weight, postnatal weight gain or adult lung function between those with and without missing covariate data, and restricting the sex and height adjusted analyses to those with full covariate data provided similar results (not shown).

A limitation of this study is our inability to adjust for other factors that may confound or mediate the association between low birth weight and reduced lung function. For example, we have little information on lower respiratory tract infections in early life. Low birth weight babies may be more susceptible to such infections and these could impair lung development. We have no information on maternal diet during pregnancy. We also have little information on childhood diet although adjusting for breastfeeding made no material difference to any of the analyses (data not shown).

Taken together, these findings lend support to the hypothesis that eventual adult lung function is partly determined by fetal growth, and to a lesser extent may be determined by infant growth. The size of these effects seems modest—based on the adjusted analyses (table 3), a 1 kg (30%) increase in birth weight is associated with increases in lung volumes and diffusing capacity of between 2% and 4%. The clinical and public health significance of these differences is uncertain. However, birth weight is, at best, only a crude measure of fetal development and hence these analyses may underestimate its influence on adult lung function. It is becoming increasingly apparent that lung function has an impact on cardiovascular and all-cause dysfunction.


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mortality as well as on respiratory outcomes. It remains to be seen whether attempts to enhance interuterine and postnatal growth can have a longlasting influence on lung development.

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