Longitudinal assessment of lung function in extremely prematurely born children

Jessica Lo MSc¹, Sanja Zivanovic PhD²,³, Alan Lunt BSc²,³, Mireia Alcazar-Paris BSc²,³, Gwendolyn Andradi MBBS²,³, Mark Thomas FRCPCH⁴, Neil Marlow FRCPCH⁵, Sandy Calvert FRCPCH⁶, Janet Peacock PhD⁷,⁸*, Anne Greenough MD²,³,⁸*

* Greenough and Peacock contributed equally to this study

¹School of Psychiatry, UNSW Medicine, University of New South Wales, Sydney, Australia
²MRC and Asthma UK Centre in Allergic Mechanisms of Asthma, King’s College London;
³Women an Children’s Health, School of Life Course Sciences, Faculty of Life Sciences and Medicine, King’s College London; ⁴Neonatal Medicine, Chelsea & Westminster Hospital, London; ⁵Neonatal Medicine, Institute for Women’s Health, London; ⁶Child Health, St. George's Hospital, London; ⁷School of Population Health and Environmental Sciences, Faculty of Life Sciences and Medicine, King's College London; ⁸NIHR Biomedical Research Centre at Guy’s and St Thomas’ NHS Foundation Trust and King’s College London.

Corresponding author: A Greenough, NICU, 4th Floor, Golden Jubilee Wing, King’s College Hospital, Denmark Hill, London SE5 8RS, UK. Tel: 0203 299 3037; Fax: 0203 299 8284; email: anne.greenough@kcl.ac.uk

Financial support: The study was funded by National Institute for Health Research (NIHR) Health Technology Assessment Programme number 08/116/10 (a detailed report of the study will be published on their website). The South London Comprehensive Local Research Network supported an administrator (RO) to facilitate recruitment. NM receives part funding from the Department of Health’s NIHR Biomedical Research Centre’s funding scheme at UCLH/UCL. The research was supported by the National Institute for Health Research
(NIHR) Biomedical Research Centre based at Guy's and St Thomas' NHS Foundation Trust and King's College London. The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the Department of Health.

**Key words**: airway function; extreme prematurity; lung volume; small airway function

**Running head**: Longitudinal assessment of lung function
ABSTRACT

Objectives: To assess longitudinally small airway function in children born extremely prematurely and whether there was a correlation between airway function in infancy and at 11 to 14 years.

Working hypotheses: There would be tracking of airways obstruction and small airway function would deteriorate during childhood in those born extremely prematurely.

Study design: A longitudinal study.

Patient – subject selection: Thirty-five children with a mean gestational age of 26 weeks had lung function assessed at one year corrected and 11-14 years of age.

Methodology: Lung volumes were measured by helium gas dilution (FRC$_{He}$) and plethysmography (FRC$_{pleth}$) and small airway function assessed by calculating the FRC$_{He}$:FRC$_{pleth}$ ratio. Airway function was assessed at one year corrected by measurement of airway resistance (R$_{aw}$) and at 11-14 years by assessment of R$_{aw}$, forced expiratory flow from 75% of vital capacity (FEF$_{75}$) and forced expiratory volume at one second (FEV$_1$).

Results: At the first assessment, the children had a mean (SD) FRC$_{He}$: FRC$_{pleth}$ of 0.90 (0.13) and at the second, 0.83 (0.12) (p= 0.035). There was a significant 0.54% decrease (95% CI: -1.02%, -0.06%) in FRC$_{He}$: FRC$_{pleth}$ for increased age per year after adjusting for birth weight, gestational age, sex and bronchopulmonary dysplasia (p=0.027). There were significant correlations between R$_{aw}$ at the first assessment and R$_{aw}$ (p=0.012), FEF$_{75}$ (p=0.034) and FEV$_1$ (p=0.04) at 11-14 years.

Conclusions: These results demonstrate in those born extremely prematurely there is tracking of airway function during childhood.
INTRODUCTION

Prematurely born infants frequently suffer troublesome respiratory symptoms and lung function abnormalities in the first two years after birth. Adolescents 1-5 and adults 6-8 born very prematurely have been reported to have airways obstruction and impaired exercise tolerance. Small airway function appears to deteriorate during infancy 9,10. It is not, however, clear whether there are changes in airway obstruction with increased age in older children, as there have been few longitudinal studies and their results conflicting. Certain studies have shown that lung function tracked with increasing age 11,12, whereas another documented airways obstruction at 7-9 years of age but not at 21 years 13. In the latter study, however the subjects were relatively mature at birth, their mean gestational age being 31.5 weeks. In contrast, amongst young people born very low birth weight (VLBW) who developed BPD, airway obstruction worsened between 8 and 18 years 14. In addition, amongst 87 extremely low birth weight (ELBW) survivors lung function significantly deteriorated between eight and twelve years (FEV1 from 83 to 77% expressed as the percent predicted for age, height and sex and FEV1/FVC from 84% to 78%); the deterioration was independent of the diagnosis of BPD 15. Whether there are changes in lung function during childhood in those born extremely prematurely and routinely exposed to antenatal corticosteroids and postnatal surfactant remains to be determined.

The United Kingdom Oscillation Study recruited infants born less than 29 weeks of gestational age 16. More than 90% of the infants were exposed to antenatal steroids and postnatal surfactant, hence their results are generalisable to the current, extremely prematurely born population. A subset had measurements at a corrected age of one year of lung volumes by plethysmography (FRCpleth) and helium gas dilution (FRCHe) and airways resistance (Raw) was also assessed 17. The FRCHe:FRCpleth ratio was calculated to assess small
airway function. The children were reassessed at 11 to 14 years of age with detailed lung function assessments. Our aim was to compare determine if there was a correlation between airway function at a corrected age of one year and at 11-14 years of age and determine if in those children, with paired FRC\textsubscript{He}: FRC\textsubscript{pleth} results, there had been deterioration in small airway function.

**PATIENTS AND METHODS**

The children included in this study had been recruited into the United Kingdom Oscillation Study (UKOS). Infants were randomised to their initial mode of ventilation (high frequency oscillatory ventilation or conventional mechanical ventilation) in the first hour after birth. The initial lung function and follow up study were approved by the South Thames Multicentre Research Ethics Committee and the South West London National Research Ethics Service Committee respectively. All parents gave informed written consent. All assessments were made in the Amanda Smith infant and paediatric respiratory laboratories at King’s College Hospital NHS Foundation Trust and performed according to guidelines from the American Thoracic Society and the European Respiratory Society. On arrival, a history was taken and the subject weighed, measured and examined. Appointments were deferred for two weeks if the subject had a respiratory tract infection.

**Assessment at one year corrected age**

The full details of the respiratory function assessments are reported elsewhere. In brief, all UKOS babies whose families lived in London were invited to attend King’s College Hospital in South London to participate in this study. Infants were sedated with 80-120 mg/kg of
chloral hydrate and monitored by pulse oximetry (Datex, Ohmeda 3800, Hatfield, UK). The testing procedure consisted of measurement of FRC by plethysmography (FRC_{pleth}) (Department of Medical Engineering, Hammersmith Hospital, London), FRC by helium dilution (FRC_{He}) (Equilibrated BioSystems, New York, USA) and airways resistance by plethysmography (R_{aw}). The results are expressed as the absolute values and as the percentage predicted for length FRC_{pleth} and FRC_{He}^{19}. The FRC_{He}: FRC_{pleth} ratio was calculated.

**Assessment at 11-14 years**

Full details of the respiratory measurements are reported elsewhere \(^{18}\). In brief, lung volumes were assessed by measuring functional residual capacity using a helium-dilution technique (FRC_{He}) and by plethysmography (FRC_{pleth}). All measurements were undertaken using a Jaeger Masterscreen PFT system (Carefusion Ltd, Basingstone, UK). Two measurements within 5\% of each other were averaged to calculate the final result \(^{20-22}\). The FRC_{He}: FRC_{pleth} ratio was calculated. Small-airway function was also assessed by measurement of forced expiratory flow at 75\% of the expired vital capacity (FEF\(_{75}\)) by spirometry. In addition, airway function was assessed by spirometric measurement of forced expiratory volume in one second (FEV\(_{1}\)), and plethysmographic assessment of R_{aw}. The absolute lung volumes are presented and all lung-function results are also expressed as the percentage predicted for height \(^{23, 24}\). Exposure to tobacco was estimated using urinary cotinine. A cotinine level of less than 10 ng per millilitre was defined as undetectable, a level of 10 to 30 ng per millilitre was considered to indicate passive smoking, and a level of more than 30 ng per millilitre was considered to suggest active smoking \(^{25}\).
Assessment of respiratory health

At age one year corrected, the infants were seen by their local paediatricians who, with the parents, completed a respiratory questionnaire. The parents were asked if their infant wheezed, had received any medicines for chest problems in the previous 12 months or had any hospital admissions for breathing difficulties. When the subjects were 11-14 years of age, their parents completed a questionnaire which included questions regarding respiratory problems prior to their children’s assessment.

Analysis

The change in lung function (FRC<sub>H</sub>: FRC<sub>pleth</sub>) over time was analysed using mixed models, which took into account the length of time between the two assessments for each child. Sex, gestational age groups (23-25 weeks, 26-28 weeks in keeping with the stratification at randomisation), birth weight and BPD status (BPD<sub>36</sub> defined as being oxygen dependent at 36 weeks PMA), were thought to be likely confounders and were adjusted for in the models. If the changes in lung function were significant for each additional year in age, the effect of sex, mode of ventilation and BPD status on the degree of change were examined. This was conducted by adding an interaction term of time and one of the aforementioned variables into the mixed model. If the term had a p-value of <0.05, the effect of the variable was considered to be statistically significant. In a sensitivity analysis, oxygen dependency at 28 days was used in the model instead of oxygen dependency at 36 weeks PMA. The relationship between airway function (R<sub>aw</sub> at a corrected age of one year and airway function (R<sub>aw</sub>, FEF<sub>75</sub> and FEV<sub>1</sub>) at 11-14 years was examined by the Pearson correlation or the Spearman correlation coefficient as appropriate. Differences in symptom status at each assessment were assessed for statistical significance using McNemar’s test. All analyses were performed using Stata v.13.0.
RESULTS

Seventy-six infants had pulmonary function assessments at one year corrected age and 44 of those children (58%) had further lung function assessment at age 11 to 14 years. Thirty-five children had results for FRC$_{He}$ and FRC$_{pleth}$ at both assessments and form this study’s cohort (Figure 1). Missing measurements at the first assessment were due to not being able to obtain technically acceptable recordings or the infant woke before the measurements were complete. Missing data at the follow-up assessment were due to the child’s inability to perform the test, results not within 10% of each other or severe airways obstruction (FEV$_1$ z score less than or equal to -3.5). The study cohort had a lower mean gestational age (p=0.04); and fewer of the mothers smoked antenatally (p=0.04) in comparison with those not included, but they were otherwise similar with respect to their baseline characteristics (Table 1). Thirteen percent of infants had wheezy attacks in the first year after birth and nine per cent had received antibiotics (table 1). When cotinine levels were assessed at age 11-14 years, 13% of the young people were exposed to passive smoking and 9% likely were active smokers (table 1).

At the first assessment, the children had a mean (SD) FRC$_{He}$: FRC$_{pleth}$ of 0.90 (0.13) and at the second assessment, 0.83 (0.12) (Table 2) (Figure 2). The lung function changes were then adjusted for several factors: sex, gestational age, birth weight and BPD status (BPD$_{36}$). This gave an adjusted mean difference between each child’s assessments of -0.0054, 95% CI -0.0102 to -0.0006, p=0.027. This difference is equivalent to a decrease of 0.54% (95% CI: -1.02%, -0.06%), in FRC$_{He}$: FRC$_{pleth}$ for each additional year in age.
In addition, we tested the model to see if the change in lung function was moderated by any of sex, BPD, mode of ventilation, smoking in pregnancy or passive smoke exposure at age 11-14. None of the interaction tests were statistically significant: sex (interaction p=0.37), BPD_{36} status (interaction p=0.69), mode of ventilation (interaction p=0.48), smoking in pregnancy (interaction p=0.07). Smoking in pregnancy did not appreciably affect the estimated mean change over time (0.54% vs 0.55% per year) and similarly passive smoke exposure was not significantly related to the change in lung function (p=0.62) and the interaction test would not converge. Similarly, the interaction test would not converge for active smoking ie was not significant, but only three children were active smokers as defined by their cotinine levels. The sensitivity analysis using BPD defined as oxygen dependency at 28 days gave similar results to those above using BPD_{36weeks}. (data not shown).

There were significant correlations between $R_{aw}$ at the first assessment and $R_{aw}$ (r=0.42, p=0.012), $FEF_{75}$ (r=-0.37, p=0.034) and $FEV_1$ (r=-0.37, p=0.04) at 11 to 14 years.

Questionnaire results of wheeze and use of chest medication were available at both assessments for 31 subjects. The number with wheeze had reduced from eight to three (p=0.008) and the number who used chest medications from sixteen to three (p<0.001). Respiratory hospital admission data were available for 29 subjects and had reduced from 12 requiring admission in the first year after birth to zero at 11-14 years (p<0.001).
DISCUSSION

We have demonstrated that airway resistance ($R_{aw}$) at one year corrected correlated with the results at follow-up of $R_{aw}$, FEF$_{75}$ and FEV$_1$. The results suggest that airway development was impaired. Indeed, at 11-14 years of age the median FEV$_1$ and FEF$_{75}$ results were at least minus one z score based on data from healthy term born children. Our results are supported by previous results. Extremely preterm infants (gestational ages < 28 weeks) or extremely low birth weight (ELBW < 1000 grams) were studied at 8 and 18 years. They had substantial airflow impairments at both age and a greater increase in small airway obstruction.

Importantly, those who were smokers at 18 years had airway obstruction that increased over time compared with those who did not. Similarly, in ELBW children the obstructive pattern of lung function deteriorated between 8 to 12 years.

A possible explanation for our results is that exposure of the immature lung to the shear stress of mechanical ventilation and/or the higher inspired oxygen concentration compared to in utero may have affected subsequent airway development. Cyclic stretching of isolated lung cells results in production of IL-8 and other inflammatory markers and the magnitude of production was related to the degree of stretch. An alternative explanation is that accelerated maturation resulting from intrauterine stress or antenatal corticosteroid administration may have adversely affected airway development. In a non-randomised study, fourteen year olds exposed to antenatal steroids compared to those unexposed had a greater prevalence of larger airway obstruction (35% versus 21%) and were twice as likely to have wheezing in the last 12 months. In our study, all of the subjects had been exposed to antenatal steroids. Such exposure, however, cannot be the only factor, as in one study reporting deterioration of small airway function during infancy, only one of the cohort had antenatal steroid exposure. A further explanation to be considered is the impact of maternal...
antenatal smoking and active adolescent smoking. Antenatal smoke exposure is associated with a reduction in airway function. Parental and active smoking have been demonstrated to act synergistically to affect early lung function deficits in young adulthood. Thirteen percent of mothers in this study smoked antenatally, but neither maternal smoking in pregnancy nor passive smoking adversely influenced the change in lung function with time. Urinary cotinine analysis indicated that 9% of the young people were likely to have been actively smoking at 11 to 14 years of age, but this amounted to only three children and no significant effect was shown in the change in lung function.

The mean FRC_{He}:FRC_{pleth} results were statistically significantly lower at 11-14 years compared to one year corrected. This result remained significant after adjusting for birth weight, gestational age, sex and BPD_{36} status. Those results suggest a deterioration in small airway function with increasing age in the extremely prematurely born young people, who had routinely been exposed to antenatal steroids and postnatal surfactant. Those results however should be interpreted with the caveat that although similar techniques were used at both assessments, different equipment was used to measure FRC_{pleth}. It has been shown that FRC measured by certain commercially available equipment was significantly lower than published reference data and lower than data reported for gas dilution techniques. We, however, used a conventional plethysmograph (Department of Medical Engineering, Hammersmith Hospital, London, UK) which has been shown to accurately measures volumes as assessed using a lung model. The FRC_{He}:FRC_{pleth} which assesses the raw data is not affected by the differences in the reference ranges. The FRC_{He}:FRC_{pleth} ratio in infancy was similar to that reported in term born infants. We did not report our Raw results as z scores as there is no one reference range which spans the age range of those included in this longitudinal study. Using a reference range which included infants and one which included
children aged 2 to 18 years \textsuperscript{38}, however, suggests that the Raw results adjusted for weight may not have changed. Hence, we suggest the change in airway function we report may be specific to small airways. The reduction in small airway function was not associated with an increase in symptoms, indeed, there was a statistically significant reduction in the respiratory hospital admission rate. In the first two years, hospitalisation is usually due to lower respiratory tract infections often due to respiratory syncytial virus \textsuperscript{39} and this becomes less of a problem with increasing age. It is possible, however, that children “live within their lung function”. In one study, lung function was highest in highly physically active children aged 9 to 10 years \textsuperscript{40}. Those results, however, could be interpreted that children with poorer lung function exercise less and, therefore, complain of less symptoms.

There are strengths and some limitations of our study. Similar techniques to assess FRC\textsubscript{He} and FRC\textsubscript{pleth} were used at both assessments, but at one year the infants were assessed supine, whereas at 11 to 14 years the children were assessed while sitting. The latter position would have “benefitted” their lung function, because in the supine position the abdominal contents would have pressed on the diaphragm reducing lung volume. Hence, we might have expected to see a higher FRC\textsubscript{He}:FRC\textsubscript{pleth} ratio, rather than the reverse. We were unable to assess all the children assessed at one year at 11 to 14 years, but those reassessed did not differ significantly from those UKOS children not included, other than this study cohort were born at a shorter mean gestational age and a lower proportion of their mothers had smoked antenatally. This suggests our results are generalisable. We did not adjust the lung function results for ethnicity, but more than 80\% of our subjects were Caucasian. A few of the children had FRC\textsubscript{He}:FRC\textsubscript{pleth} ratios that were greater than one (Figure 2), but as each of their lung volume results fulfilled the ATS guidelines for reproducibility their data were included in the analysis. We did not report our Raw results as percentage predicted for height as our
Ethics Committee did not allow us to sedate healthy term born infants for the purpose of creating a reference range. In this subset, only 10% had wheeze at 11-14 years compared to 15% in the whole cohort, suggesting our sample was not biased by parents seeking further assessment for a symptomatic child.

In conclusion, we have demonstrated that tracking occurs in lung function between infancy and 11 to 14 years. Our results suggest that there may have been a deterioration in small airway function. Lung function abnormalities in school age children may have important functional consequences. In one study of 9 to 11 year old children, in those born prior to 33 weeks of gestation, airways obstruction was associated with chest CT abnormalities including increased subpleural opacities, bronchial wall thickening and hypo-attenuated lung areas.
ACKNOWLEDGEMENTS

**Competing interests:** AG has held grants from various manufacturers (Abbot Laboratories, MedImmune) and ventilator manufacturers (SLE). AG has received honoraria for giving lectures and advising various manufacturers (Abbot Laboratories, MedImmune) and ventilator manufacturers (SLE). AG is currently receiving a non conditional educational grant from SLE.

**Contributor statement:** AG and JP designed the study. SZ, MAP, AL, GA and MT collected the data. JL and JP analysed the data. All of the authors critically reviewed the manuscript and approved the final manuscript as submitted.

JLP is a NIHR Senior Investigator.
REFERENCES


FIGURE LEGENDS

Figure 1: Flow chart for study cohort

Figure 2: $\text{FRC}_\text{He}:\text{FRC}_\text{pleth}$ results at the first and second assessment

TABLE LEGENDS

Table 1: Baseline characteristics by inclusion status

Table 2: Lung function results (unadjusted) at the initial and second assessment