Progressive static pulmonary hyperinflation in survivors of severe bronchopulmonary dysplasia by mid-adulthood

ABSTRACT

Background: Severe bronchopulmonary dysplasia (BPD) might be associated with an accelerated age–related decline of lung function.

Methods: 14 individuals were studied longitudinally at 15±4, 18±3 and 38±3.2 years. Information on personal history was completed, and lung function testing and skin prick testing were performed. Longitudinal data were compared intra-individually and with matched controls from the NHANES III dataset.

Results: The ratio of residual volume/total lung capacity (RV/TLC) increased markedly from 25.9±7.0% to 39.3±6.8%. A significant time-effect was found compared to controls for the forced vital capacity (FVC) which decreased more rapidly than expected. Flow values were at the lower limit of normal range but remained relatively stable over time. Some individuals had completely normal lung function results.

Conclusion: Increasing static pulmonary hyperinflation with age is indicative of bronchiolar dysfunction or early emphysematous changes in survivors of severe BPD. Susceptibility for long-term sequelae shows significant variability.

In 1967, Northway et al reported on harmful effects of barotrauma and oxygen toxicity in prematurely born children, causing an acquired lung disease called bronchopulmonary dysplasia (BPD) that was characterised by patchy bronchoalveolar scarring, atelectases and overexpansion. With the introduction of lung protective ventilation and the advent of surfactant, the features of BPD have changed. Residual lung function abnormalities, mainly airflow obstruction and hyperinflation, were found in young adults, and emphysematous changes found by CT raised the question whether BPD might ultimately affect pulmonary ageing, for example, by leading to the development of chronic obstructive pulmonary disease.

A unique cohort of 20 survivors of severe BPD born before the advent of lung protective ventilation and surfactant has been observed longitudinally into adulthood. Neonatal pulmonary management had included pressure-controlled mechanical ventilation with peak inspiratory pressures up to 40 cm H2O adjusted to repeated blood gas analyses; FiO2 values of 0.3, 0.4, 0.75 or 1.0 and 1 to 2 hourly suctioning and at least 48-hourly changing of the endotracheal tubes. The individuals had undergone pulmonary function testing at 15±4 years (1983) and 18±3 years (1987). Sixteen of them could be contacted for follow-up at 38±3.2 years, and 14 (5 women) consented to participate. Subjects underwent skin prick testing to prevalent aeroallergens and pulmonary function testing. Each individual was matched for age, sex and height with one to four individuals from the National Health and Nutrition Examination Survey III dataset, both for the 1987 and the 2008 follow-up (courtesy of Dr J.L. Hankinson, Valdosta, GA, USA). Further details are outlined in the online supplement.

The perinatal characteristics were birthweight 1795±456 g (mean±SD), gestational age 31.8±2.9 weeks, highest documented peak inspiratory pressures 26±4.5 cmH2O, endotracheal intubation for 38 (range 4–105) days and FiO2 ≥0.4 for 36 (range 4–96) days. Five individuals had been tracheotomised for an extended period. The first sequential lung function study in 1983 and 1987 had revealed mean flow parameters around the lower limit of normal, but no significant changes over time.

Six individuals reported a variety of persisting respiratory symptoms, mainly asthma, cough or shortness of breath during exercise. Four individuals with hay fever were sensitised to pollens; an additional person was sensitised to cat allergens without symptoms. Two individuals were on occasional inhaled corticosteroids. There were four current smokers (8–16 pack-years) and one former smoker. Six individuals had shown bronchial hyperreactivity to methacholine in 1987, two of whom still reacted.

Table 1 Lung function data of the 14 survivors of BPD participating in both follow-up studies and shift of mean differences between patients and controls from 1987 to 2008

<table>
<thead>
<tr>
<th>Shift of mean difference 1987–2008</th>
<th>95% CI</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>FVC (L)</td>
<td>−0.584</td>
<td>−1.092 to −0.076</td>
</tr>
<tr>
<td>FEV1 (L/sec)</td>
<td>−0.367</td>
<td>−0.971 to 0.237</td>
</tr>
<tr>
<td>FEV1/FVC</td>
<td>0.003</td>
<td>−0.138 to 0.143</td>
</tr>
</tbody>
</table>

Numbers indicate means±SD.

FEV1, forced expiratory volume in the first second; FRC, functional residual capacity; FVC, forced vital capacity; MEF25, 50, maximum expiratory flows at 25% and 50% of vital capacity; RV, residual volume; TLC, total lung capacity.
in 2008. One originally negative person showed moderate reactivity in 2008. Transfer factor of the lung for carbon monoxide in 2008 was 83.5±15.7% (mean±SD, range 58–107%) of predicted; FeNO was 10.8±6.5 (range 4–27 ppb). The results of pulmonary function testing are depicted in table 1 and figure 1. Smoking and other potential confounders had no significant effect; lowest flow values were found among non-smokers.

The results are consistent with a progressive static hyperinflation into mid-adulthood, suggestive of progressive bronchiolar dysfunction or early emphysematous changes with ageing in severe BPD. The relatively minor flow changes during the 20-year period indicate dysfunction of small airways and are dissimilar to chronic obstructive pulmonary disease, hinting to differing disease mechanisms. The current study is in line with the concept of adult lung disease originating early in life. Susceptibility for long-term sequelae shows significant variability.

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Competing interests None.

Ethics approval This study was conducted with the approval of the Ethics Committee of Basel (ref. no. 338/07).

Contributors DT was primary investigator in all aspects of the study. MHB significantly contributed to the design of the study and interpretation of results from the adult pulmonologist’s perspective. HH-B was primary investigator of the first follow-up study and added to the manuscript. JH helped with the design, the execution and the analysis of the study and writing the manuscript.

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REFERENCES


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