ORIGINAL ARTICLE

The association between childhood asthma and adult chronic obstructive pulmonary disease

Andrew Tai,1 Haily Tran,2 Mary Roberts,3 Nadeene Clarke,2 John Wilson,4 Colin F Robertson2,5

ABSTRACT

Introduction There is epidemiological evidence to suggest that events in childhood influence lung growth and constitute a significant risk for adult COPD. The aim of the study is to evaluate for an association between childhood asthma and adult COPD.

Methods This longitudinal, prospective study of 6–7-year-old children with asthma has been regularly reviewed every 7 years to the current analysis at 50 years of age. Participants completed respiratory questionnaires and lung function spirometry with postbronchodilator response. At the age of 50, subjects were classified to the following subgroups: non-asthmatics, asthma remission, current asthma and COPD which was defined by FEV1 to FVC ratio postbronchodilator of less than 0.7.

Results Of the remaining survivors, 346 participated in the current study (participation rate of 76%) of whom 197 completed both questionnaire and lung function testing. As compared with children without symptoms of wheeze to the age of 7, (non-asthmatics) children with severe asthma had an adjusted 32 times higher risk for developing COPD (95% CI 3.4 to 269). In this cohort, 43% of the COPD group had never smoked. There was no evidence of a difference in the rate of decline in FEV1 (mL/year, 95th CI) between the COPD group (17, 10 to 23) and the other groups: non-asthmatics (16, 12 to 21), asthma remission (20, 16 to 24) and current asthma (19, 13 to 25).

Conclusions Children with severe asthma are at increased risk of developing COPD.

INTRODUCTION

COPD is an umbrella term for a spectrum of diseases characterised by airflow obstruction that is not fully reversible, resulting from a combination of small airway disease and parenchymal destruction. While the main risk factors for the development of COPD are genetic and inhalational exposures such as tobacco smoke, failure to attain maximal lung growth during childhood constitutes a significant risk. In the 1970s, Burrows et al1 reported that adults with COPD were more likely to recall respiratory troubles during childhood than those without COPD, suggesting the possible progression of childhood asthma into adult COPD. In the early 1960s, Orie and colleagues proposed that “asthma, chronic bronchitis and emphysema should be considered different expressions of one disease entity in which both host factors (such as genetic predisposition to atopy and bronchial hyperresponsiveness) and environmental factors (such as smoking) play major roles”.2

Key messages

What is the key question?
▸ Do children with asthma develop COPD in adult life?

What is the bottom line?
▸ Children with severe asthma are at increased risk of developing adult COPD whereas children with intermittent asthma are not at increased risk of developing adult COPD.

Why read on?
▸ This study is the longest and most comprehensive study of childhood asthma followed to the age of 50 and highlights that suboptimal lung growth in childhood is a precursor to adult COPD.

In the Tuscon Epidemiological Study of Airways Obstructive Disease (TESAOD), 3099 adult subjects over 20 years of age were randomly selected and subsequently followed up over 20 years.3 The results of the study indicated that adult subjects with active asthma had a 10 times higher risk of acquiring symptoms of chronic bronchitis, 17 times higher risk of receiving a diagnosis of emphysema and 12.5 times higher risk of fulfilling COPD criteria, even after adjusting for smoking history and potential confounders. In the large follow-up of the European Community Respiratory Health Survey, subjects with a history of childhood asthma had a 10-fold increased risk of developing COPD.4 In the most recent follow-up of the Tuscon birth cohort to the age of 22, poor airway function shortly after birth was recognised as a risk factor for airflow obstruction in young adults and these subjects would reach the threshold of FEV1 and FEV1/FVC ratio that define COPD at an earlier age.5

The synergistic impact of smoking and asthma on lung function decline has been demonstrated in adult studies.6 7 with the Dunedin cohort reporting follow-up from childhood years and demonstrating that smoking was a risk factor for persistent wheezing and subsequent reduced lung function at the age of 26.6

It is increasingly apparent that COPD often has its roots decades before the onset of symptoms. Impaired growth of lung function during childhood and adolescence, caused by premature gestation, asthma, recurrent infections or tobacco smoking,
may lead to lower maximally attained lung function in early adulthood and also predispose to development of COPD.9 Therefore, the aim of this study was to explore the association between childhood asthma into adult COPD. Some of the results of this study have been previously reported in the form of an abstract.10

METHODOLOGY
This study was approved by the Human Research Ethics Committee of the Royal Children’s Hospital Melbourne. All subjects provided consent prior to participation.

Study design and cohort
The Melbourne Asthma Cohort was recruited from a 1957 birth cohort at the age of 7 which has been previously described11 and comprehensively reviewed from the age of 7 to the current study at the age of 50.12–16 In brief, a modified random sampling strategy was used to select 401 subjects, following a survey of 30 000 Grade 2 Melbourne primary school children in 1963–1964. The survey included a parent completed questionnaire which was followed by a parent interview and physical examination of the child by members of the research team. All those children with a history of asthma or wheezy bronchitis (WB) were included in the study and one in two with mild wheezy bronchitis (MWB) and one in 20 controls were included. When the children were reviewed at the age of 10, it was realised that there were very few with severe asthma (SA), and a further sampling of 21 000 children was performed from the same birth cohort to establish an SA group which included 83 children.

The original groups were classified as follows: control or non-asthmatics—105 children who had never wheezed; MWB—74 children with fewer than five episodes of wheezing associated with bronchitis; WB—104 children with five or more episodes of wheezing associated with bronchitis; asthma—113 children with wheezing not associated with respiratory tract infection; and SA—83 children with onset of symptoms before 3 years of age, persistent symptoms at 10 years of age, barrel-chest deformity and or FEV1 to FVC ratio to 50% or less (of which only three had a FEV1 to FVC ratio under 50%). The group of children with MWB and WB would now be regarded as children with intermittent asthma.

Interviewer-administered questionnaire
At the current study, an interviewer-administered questionnaire was performed to collect details on the frequency of wheeze, symptoms of chronic bronchitis (defined as daily moist productive cough for at least 3 months over two successive years), frequency of hay fever and eczema symptoms and smoking behaviour. Childhood hay fever (data collected at the age of 7) was defined as children who had symptoms of sneezing, nasal itching, runny or blocked nose in the absence of a cold or flu at recruitment. Childhood eczema (data collected at the age of 7) was defined as children who had symptoms of dry, itchy rash localised to flexural regions, facial or generalised to the body at recruitment. Current smoker was defined as subjects who smoked regularly within the last 12 months. Ever smoker was defined as subjects who had a smoking history of at least 1 pack year or more. The St George’s Respiratory Questionnaire was used as a valid measure of activity limitation with a low score indicating a better health related quality of life. The threshold for a clinically significant difference between and within groups of patients is four units.17 Subjects who were not able to present to the hospital had the questionnaire administered by telephone.

Lung function testing
Lung function testing was performed on all subjects who presented to the hospital for assessment. The test was carried out by a trained respiratory scientist using the Masterscreen body plethysmograph (Jaeger, Germany). Forced expiratory manoeuvres were performed according to American Thoracic Society/European Respiratory Society standards.18 FEV1, FVC and FEV1 to FVC ratio were the variables used for analysis. Spirometry was repeated following administration of 600 μg dose of salbutamol. A postbronchodilator increase in FEV1 of 200 mL or 12% defined reversibility.18 COPD was defined according to the GOLD criteria: FEV1 to FVC ratio postbronchodilator of less than 0.7.19

Statistical analysis
For the follow-up at 50 years, the cohort was classified as follows.

- Non-asthmatics: Those who were originally in the non-asthmatic childhood group and remained wheeze-free throughout the study period.
- Asthma remission: Those who were originally in one of the four symptomatic groups at recruitment who had been wheeze-free and not taken asthma medication in the preceding 3 years.
- Current asthma: Those who were originally in one of the four symptomatic groups or non-asthmatic group at recruitment and who had asthma symptoms or had taken asthma medication for the preceding 3 years.
- COPD: Postbronchodilator FEV1 to FVC ratio <0.7 group who were defined at the age of 50.

The statistical software Stata V10.0 for Windows (Stata Corporation, Texas, USA) was used for data analysis. Symptom outcomes of the age 50 groups were determined with χ2 testing compared with the non-asthmatic group. Univariate and multivariate logistic regression was used to determine the clinical predictors of childhood factors into adult COPD. The childhood non-asthmatic group was used as the reference comparison group. The following variables were tested on multivariate analysis: childhood SA, asthma, WB groups, male sex, ever smoker, current smoker at the age of 50, childhood hay fever and childhood eczema. Two-sided Student t tests were used to compare prebronchodilator and postbronchodilator lung function measures at the age of 50 between the age 50 groups and the non-asthmatic group. The rate of decline in FEV1 was calculated separately for each subject with at least two time point measurements between the ages of 21 and 50 via a linear regression. The age of 21 was taken as the time when maximal lung function would have been attained. The effects of smoking at the age of 21 (binary outcome), smoking at the age of 50 (binary outcome), and sex on the rate of decline in FEV1 were examined using multivariate linear regression. The rate of decline in FEV1 to FVC ratio was calculated separately for each subject with at least two time point measurements between the ages of 10 and 50 via a linear regression. The overall difference in the rate of decline in FEV1 and FEV1 to FVC ratio across the age 50 groups was examined using an analysis of variance.

RESULTS

Subject recruitment
Of the original 484 participants, 21 had died (2 due to asthma) and five original records were lost. Of the available 458 participants, 34 refused contact and 78 were lost to follow-up. In all,
42 subjects with COPD and chronic bronchitis at the age of 50.

COPD demographics
In this cohort, 28/197 (14%) subjects had COPD. Of the 28 subjects, 22 (79%) were men and 6 (21%) were women. In the COPD group, 16/28 (57%) had a history of smoking and 12/28 (43%) had never smoked. The prevalence rates of both current smokers and ever smokers were proportional in all groups at 50 years of age.

Outcomes at the age of 50
Children in the MWB and WB groups had the highest prevalence of asthma remission at the age of 50 (48% and 56%, respectively) compared with the SA group (15%). Overall, 44% of children with SA fulfilled the diagnosis of COPD at the age of 50 (table 2). In this study, 7/28 (25%, p=0.01) of the subjects with COPD at the age of 50 had symptoms of chronic bronchitis compared with 1/41 (2%) of the non-asthmatic, 2/58 (3%, p=0.8) of asthmatic remission and 11/70 (16%, p=0.03) of asthma remission and 11/70 (16%, p=0.03) of current asthma groups during the age of 50 years.

The childhood predictors of adult COPD
On univariate analysis, a significantly increased OR (OR, 95th CI) was found for the childhood asthma and SA (37.1, 4.6 to 301) groups and the diagnosis of COPD at the age of 50. Following multivariate analysis, the increased OR was strengthened for the childhood SA group. There was no significantly increased OR for sex, childhood WB groups, childhood hay fever or eczema, current smoker or ever smoker categories with COPD at 50 years of age (table 3).

Lung function at the age of 50 and trend from childhood years
At the age of 50, the mean FEV₁% predicted (95th CI) in those with current asthma (92, 88 to 97; p<0.005), asthma remission (104, 100 to 108; p<0.05) and COPD (85, 81 to 90; p<0.005) were reduced compared with those who were in the non-asthmatic group (111, 107 to 115). Mean FEV₁ to FVC ratio (95th CI) in those with current asthma (71, 69 to 73, p<0.005) and COPD (63, 61 to 65; p<0.005) were reduced compared with those who were in the non-asthmatic group (78, 76 to 80).

Further analysis of the group with COPD demonstrated that the mean FEV₁%predicted (95th CI) in those who were smokers (86, 78 to 94; p=0.7) were similar to those who were non-smokers (85, 79 to 90). The mean FEV₁ to FVC ratio (95th CI) in those who were smokers (63, 60 to 66; p=0.8) were similar to those who were non-smokers (63, 60 to 65).

The reduced lung function seen in those with COPD and current asthma at 50 years was clearly established in the childhood years (figure 1). In the COPD group, there is a lack of increase in %predicted FEV₁ when compared with the non-asthmatic, asthma remission and current asthma groups during the age of 14–21.

The rate of decline in lung function across recruitment of age 50 groups
For each group at age 50, the rate of decline and 95% CI in FEV₁ (mL/year) from age 21 to 50 and FEV₁ to FVC ratio (percentage points/year) from age 10 to 50 is shown in figure 2. The rate of decline in FEV₁ (mL/year, 95th CI) is similar across all groups: non-asthmatic 16 (95th CI 12 to 21), asthma remission 20 (95th CI 6 to 24), current asthma 19 (95th CI 13 to 25) and COPD 17 (95th CI 10 to 23) There was no evidence of an association among the rate of decline in FEV₁ and sex, smoking at the age of 21 and smoking at the age of 50.

Table 1 Distribution of subjects at recruitment and the age of 50

<table>
<thead>
<tr>
<th>Recruitment</th>
<th>Deceased</th>
<th>Followed up</th>
<th>Lung function test performed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age 50</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control (non-asthmatics)</td>
<td>105 (22%)</td>
<td>5</td>
<td>77 (22%)</td>
</tr>
<tr>
<td>Mild wheezy bronchitis</td>
<td>74 (15%)</td>
<td>3</td>
<td>50 (14%)</td>
</tr>
<tr>
<td>Wheezy bronchitis</td>
<td>104 (22%)</td>
<td>6</td>
<td>78 (23%)</td>
</tr>
<tr>
<td>Asthma</td>
<td>113 (24%)</td>
<td>2</td>
<td>81 (24%)</td>
</tr>
<tr>
<td>Severe asthma</td>
<td>83 (17%)</td>
<td>5</td>
<td>60 (17%)</td>
</tr>
<tr>
<td>Male (%)</td>
<td>61%</td>
<td>81%</td>
<td>60%</td>
</tr>
</tbody>
</table>

Note that five original records were lost, altering the recruitment total to 479 subjects and 21 subjects were deceased, leaving 458 survivors. The percentages followed up are based on the survivors.

Table 2 Outcome at the age of 50 by recruitment groups

<table>
<thead>
<tr>
<th>Recruitment</th>
<th>Asthma remission (n=57)</th>
<th>Current asthma (n=65)</th>
<th>COPD (n=27)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MWB</td>
<td>11/23 (48%)</td>
<td>11/23 (48%)</td>
<td>1/23 (4%)</td>
</tr>
<tr>
<td>WB</td>
<td>24/43 (56%)</td>
<td>16/43 (37%)</td>
<td>3/43 (7%)</td>
</tr>
<tr>
<td>A</td>
<td>17/49 (35%)</td>
<td>24/49 (49%)</td>
<td>8/49 (16%)</td>
</tr>
<tr>
<td>SA</td>
<td>5/34 (15%)</td>
<td>14/34 (41%)</td>
<td>15/34 (44%)</td>
</tr>
</tbody>
</table>

1/48 of the control group had COPD at the age of 50.
A, asthma; MWB, mild wheezy bronchitis; SA, severe asthma; WB, wheezy bronchitis.

Table 3 The childhood predictors of adult COPD

<table>
<thead>
<tr>
<th>OR (95% CI)</th>
<th>OR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(univariate)</td>
<td>(multivariate)</td>
</tr>
<tr>
<td>Severe asthma</td>
<td>37.1 (4.6 to 301)</td>
</tr>
<tr>
<td>Asthma</td>
<td>9.1 (1.1 to 76.4)</td>
</tr>
<tr>
<td>Wheezy bronchitis</td>
<td>3.5 (0.4 to 35.2)</td>
</tr>
<tr>
<td>Mild wheezy bronchitis</td>
<td>2.1 (0.1 to 35.8)</td>
</tr>
<tr>
<td>Male sex</td>
<td>2.4 (0.9 to 6.3)</td>
</tr>
<tr>
<td>Ever smoker</td>
<td>1.0 (0.5 to 2.3)</td>
</tr>
<tr>
<td>Current smoker</td>
<td>1.1 (0.5 to 2.4)</td>
</tr>
<tr>
<td>Childhood hay fever</td>
<td>1.0 (0.3 to 3.8)</td>
</tr>
</tbody>
</table>
There was no evidence of a difference in the rate of decline in FEV$_1$ to FVC ratio (% points/year, 95th CI) across the age 50 groups: non-asthmatic 0.27 (95th CI 0.22 to 0.32); asthma remission 0.25 (95th CI 0.21 to 0.29); current asthma 0.27 (95th CI 0.21 to 0.32); and COPD groups 0.27 (95th CI 0.18 to 0.37).

CONCLUSIONS
In this longitudinal follow-up study, it is clear that children with SA are at increased risk of developing adult COPD by 50 years of age. Children with intermittent asthma and persistent asthma are not at increased risk of developing COPD. When compared with the recent follow-up of the European Community Respiratory Health Survey, subjects who had reported a childhood history of asthma similarly reported a 10-fold increased risk of developing COPD.4

Findings from cross-sectional studies of hospital-based samples of asthmatics suggest that asthma alone can cause persistent, non-reversible airflow obstruction and that the degree of obstruction is a function of the duration and severity of previous asthma.20 21 Reversibility of pulmonary obstruction in response to treatment, a hallmark of asthma, may decrease over time in some patients with moderate or SA, to the point of irreversible or only partially reversible airway obstruction. In the TESAO study group, the investigators did not have information on response to bronchodilator, leaving unanswered the question of progression of asthma into COPD.3

The lung function outcome of the group with COPD adds to the work of Stern et al3 who followed the Tucson birth cohort group indicating that infants with reduced lung function continue to track at lower levels to the age of 22. The results of this study highlight the importance of suboptimal lung function growth in childhood as a predisposing factor towards adult COPD. Recent reviews by Bush22 and Narang and Bush23 highlight the interaction between adverse maternal factors (such as smoking, atopy, infection) on the antenatal lung, childhood factors (such as candidate genes, asthma, infections, premature gestation) on lung growth and environmental risk factors (such as air pollution, smoking) as predisposing factors towards the development of COPD. During the period of adolescence, it was also noted that the adults who subsequently developed COPD did not have an increase in lung function growth as compared with the other groups which may suggest another mechanism contributing as a precursor to the development of COPD.

Tobacco smoke is by far the most important risk factor for COPD worldwide, but it is well recognised that only a fraction of smokers develop COPD and no less than a fourth of cases of COPD are unrelated to smoking.24 In the past decade, there have been emerging population-based studies reporting non-smoking COPD as much higher than previously believed with prevalence reported between 15% and 50% worldwide. Other risk factors included air pollution, occupational exposures, lower respiratory childhood infections, chronic asthma, poor nutrition and environment interaction with candidate genes.25

As part of the Copenhagen City Heart Study, 9% of subjects who were non-smokers developed COPD over a 25-year follow-up.26 In this cohort, a substantial subset of our COPD cohort have never smoked, confirming a phenotype of COPD.
which is potentially related to childhood asthma. In our cohort, 25% of the subjects with COPD had symptoms of chronic bronchitis, adding to the heterogeneous nature and varying phenotype of COPD in adult life. Beasley and colleagues suggest that COPD should be considered as a syndrome of different disorders with distinct pathophysiological processes and treatment responses and, therefore, it would be important to define, identify and treat the different components of this complex disorder. The synergistic impact of smoking and asthma on lung function has been demonstrated in the adult studies as well.6 9 This has not been well described in longitudinal childhood asthma studies. The combination of asthma and smoking >15 cigarettes per day had a synergistic effect on the decline in lung function, and resulted in a 17.8% decline in FEV1 over 10 years in the CARDIA study.7 In this study, we found no increase in the rate of decline in lung function of the current asthma and COPD groups when compared with the non-asthmatics when the smoking behaviour was taken into account. Due to the difficulty in extracting accurate smoking behaviour and varying subject attendance at different time points, we were not able to subdivide the impact of smoking behaviours (eg, continuous smokers, quitters) and total pack year smoking history on the decline in lung function. Therefore, the true impact of smoking has to be interpreted with caution as it is clearly underpowered to measure the effect on lung function decline.

Airway remodelling is characterised by functionally relevant changes of the cellular and molecular constituents of the airway wall that are linked to the surrounding inflammatory process.28 29 Airway inflammation and remodelling have been documented in young children with asthma30–32 and is postulated to be linked among patients with asthma to the development of non-reversible airway obstruction and accelerated FEV1 decline.33–35 However, to date, this association is far from being fully elucidated. Indirect evidence is provided by Benayoun et al17 in which fibroblast accumulation and airway smooth muscle hypertrophy in proximal airways were found to characterise severe persistent asthma and to correlate negatively with both prebronchodilator and postbronchodilator FEV1 values among patients with asthma. An inclusion criterion for children recruited to the SA group was either FEV1 to FVC ratio <50% or barrel-chest deformity, a level of severity which is rarely seen today. These children with SA may not be representative of current children with SA as subjects in this cohort were also managed at a time when corticosteroids were not available in clinical practice until they were in their late 20s or 30s. As to whether inhaled corticosteroids have any impact on lung function of asthmatic subjects tracking into adult life, this has been addressed in the Dunedin cohort whereby subjects who were receiving inhaled corticosteroids had lower lung function at the ages of 18 and 26 compared with those not receiving inhaled corticosteroids, although this outcome may have been confounded by the severity of the underlying asthma.8 37 There are several limitations in our study including a reduced participation rate which may have led to selection bias. Despite this, we believe that our current study group is representative of the recruitment group. Data on questionnaires are also subject to recall errors. The definition of the asthma group was based on questionnaire data alone which in itself is prone to recall bias and possible error. Other risk factors such as candidate genes, prematurity, parental history of asthma, childhood respiratory infections, maternal smoking, occupational exposures to dusts and chemicals or biomass fuels are associated with the development of COPD but in this longitudinal study, data on these risk factors were not available. The small number of subjects who fulfilled the diagnosis of COPD is also a limitation and may affect the power of negative findings. Last, the absence of bronchial hyper-responsiveness testing in children does not allow measurement of its contribution as a risk factor towards adult COPD and on the decline in lung function.

In conclusion, children with SA are at increased risk of developing adult COPD, despite the absence of a smoking history. Children with intermittent and persistent asthma which accounts for the majority of childhood asthma are not at increased risk. The fixed abnormalities in lung function in adult life are clearly established in childhood and track at lower values progressing to irreversible airway obstruction in adulthood.

Correction notice This article has been corrected since it was published Online First. The author Nadeene Clark has been amended to read Nadeene Clarke.

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Contributors Acquisition of data: AT, MR, HT, NC. Analysis and interpretation of data: AT, JW, CR. Drafting or revising the paper for important intellectual content and final approval of data: AT, MR, HT, NC, JW, CR.

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REFERENCES
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