Prediction of the rate of decline in FEV₁ in smokers using quantitative computed tomography

R Yuan,1,2 J C Hogg,1,3 P D Paré,1,4 D D Sin,1,4 J C Wong,1 Y Nakano,5 A M McWilliams,4,6 S Lam,4,6 H O Coxson 1,2

ABSTRACT

Background: A study was undertaken to determine if quantitative CT estimates of lung parenchymal over-inflation and airway dimensions in smokers with a normal forced expiratory volume in 1 s (FEV₁) can predict the rapid decline in FEV₁ that leads to chronic obstructive pulmonary disease (COPD).

Methods: Study participants (n = 143; age 45–72 years; 54% male) were part of a lung cancer screening trial, had a smoking history of >30 pack years and a normal FEV₁ and FEV₁/forced vital capacity (FVC) at baseline (mean (SD) FEV₁ 99.4 (12.8) %, range 80.2–140.7%; mean (SD) FEV₁/FVC 77.9 (4.4), range 70.0–88.0%). An inspiratory multislice CT scan was acquired for each subject at baseline. Custom software was used to measure airway lumen and wall dimensions; the percentage of the lung inflated beyond a predicted maximal lung inflation, the low attenuation lung area with an x-ray attenuation lower than −950 HU and the size distribution of the overinflated lung areas and the low attenuation area were described using a cluster analysis. Multiple regression analysis was used to test the hypothesis that these CT measurements combined with other baseline characteristics might identify those who would develop an excessive annual decline in FEV₁.

Results: The mean (SD) annual change in FEV₁ was −2.3 (4.7)% predicted (range −23.0% to +8.3%). Multiple regression analysis revealed that the annual change in FEV₁ %predicted was significantly associated with baseline percentage overinflated lung area measured on quantitative CT, FEV₁ %predicted, FEV₁/FVC and gender.

Conclusion: Quantitative CT scan evidence of over-inflation of the lung predicts a rapid annual decline in FEV₁ in smokers with normal FEV₁.

Chronic obstructive pulmonary disease (COPD) is an inflammatory lung disease caused by the inhalation of toxic particles and gases that results in destruction of the lung parenchyma and remodelling of the small airways.1 Tobacco smoking is the most important risk factor for COPD, but the fact that only a minority of smokers develop COPD strongly suggests that the host response is equally important in the pathogenesis of this condition.2,3 That only a susceptible minority of smokers develop COPD was discovered in a classic study of the natural history of chronic bronchitis and emphysema by Fletcher et al.4 This study showed that, over 8 years of follow-up, only 13% of participants experienced a decline in forced expiratory volume in 1 s (FEV₁) and therefore ended with a final FEV₁ that was low enough to satisfy the current diagnostic criteria for COPD.5 Although recent data suggest that this small fraction may have been an underestimate, the concept that only a minority of heavy smokers develop COPD has not been challenged.6,7 By the early 1970s it was recognised that the airflow limitation that defines COPD is caused by a combination of increased resistance in the small conducting airways and decreased parenchymal elasticity caused by emphysematous destruction.8,9 Although many tests have been designed to detect small airway abnormalities at an early and hopefully reversible stage, they have been largely abandoned because they failed to identify the minority of smokers with normal expiratory flows who go on to develop COPD.6,7 The introduction of non-invasive quantitative imaging of both emphysematous lung destruction and airway remodelling has provided a fresh approach to detecting changes in the anatomy of the peripheral lung. Using these imaging approaches, investigators have shown that subjects with normal lung function may have emphysematous destruction in their lungs.8,9 These observations led to the hypothesis that early emphysematous destruction might be associated with a subsequent rapid decline in FEV₁ that leads to COPD. The present study used computed tomography (CT) scans from subjects participating in a lung cancer screening study to quantify the inflation of the lung parenchyma, lung area with a lower x-ray attenuation and airway dimensions, and correlated these measurements with serial spirometry that establish a subject’s individual decline in FEV₁.

METHODS

Subjects

Subjects in the current study were from the British Columbia (BC) Cancer Agency lung cancer screening programme, the BC-Lung Health Cohort.10 This sub-cohort is composed of smokers who had normal spirometry at baseline (ie, FEV₁ ≥80% of predicted value; ratio of FEV₁ to forced vital capacity (FEV₁/FVC) ≥70%); at least two spirometry measurements at least 6 months apart; and a baseline CT scan obtained using either a GE (GE Medical System, Milwaukee, Wisconsin, USA) or Siemens scanner (Siemens Medical Solutions; Erlangen, Germany).

Lung function

Spirometry was performed using American Thoracic Society guidelines without the administration of a bronchodilator.11 FEV₁ was expressed as a percentage of the predicted value

FVC was calculated using Crapo’s equations. FEV\(_1\)/FVC was calculated using actual values. The annual change in \(\Delta\text{FEV}_{1}\%\text{predicted/\text{year}}\) was calculated for subjects with two visits as: \(\Delta\text{FEV}_{1}\%\text{predicted at T1} - \Delta\text{FEV}_{1}\%\text{predicted at T0}/\text{follow-up years}\). For subjects with more than two visits, \(\Delta\text{FEV}_{1}\%\text{predicted/year}\) was the slope of the regression line, in which all the available FEV\(_1\)%predicted measurements were plotted against age. A negative value of \(\Delta\text{FEV}_{1}\%\text{predicted/year}\) indicates worsening of the lung function.

CT technique

All CT scans were acquired in the volume scan mode at suspended full inspiration with the subject in the supine position. No intravenous contrast media were used. These CT scans were acquired using a GE scanner (Lightspeed Ultra, 120 kVp, 160 mAs, 1.25 mm slice thickness, standard reconstruction kernel) in 36 cases (25%) and using a Siemens scanner (Sensation 16, 120 kVp, 125 mAs, 1.0 mm slice thickness, BSSf reconstruction kernel) in 107 cases (75%). These two image acquisition protocols have been shown to provide comparable CT densitometry measurements.

Quantitative CT analysis

A quantitative analysis of the lung parenchyma was performed using custom software (EmphybJ) as previously described. Briefly, the lung parenchyma was segmented from the chest wall and large central blood vessels in all CT images using a modified border tracing algorithm with a prior position knowledge algorithm. Total lung volume was calculated by summing the segmented pixel area in each slice and multiplying by the slice thickness. For each pixel, the mean CT attenuation (in Hounsfield Units, HU) was calculated and converted to density by adding 1000 to the HU number and dividing by 1000, and the lung inflation (ie, volume of gas/g of tissue) was calculated according to equation 1:

\[
m\text{(gas)} = \frac{\text{Specific volume (tissue & gas)} \times \text{Specific volume (tissue)}}{g\text{(tissue)}}
\]

where specific volume is the inverse of density. The density of the lung (tissue and gas) was measured from the CT scan, and the density of gas-free tissue was assumed to be 1.065 g/ml and constant for all subjects.

The predicted total lung capacity (TLC) was obtained using the following equations (equations 2a and 2b) from Crapo et al.

Women: Predicted TLC (ml) = 59 × height (cm) − 4537 (2a)

Men: Predicted TLC (ml) = 79.5 × height (cm) + 3.2 × age (years) − 7333 (2b)

The predicted lung weight was obtained by first calculating the predicted body weight using an equation described by Devine (equations 3a and 3b), and then substituting body weight into the prediction equation for lung weight modified from that originally provided by Shohl (see online supplement) (equation 4).

Women: Predicted body weight (g) = 45500 + 905.5 × (height (cm) − 152.4) (3a)

Men: Predicted body weight (g) = 50000 + 905.5 × (height (cm) − 152.4) (3b)

Predicted lung weight (g) = 0.017 × ideal body weight (g) − 83.12 (4)

Since TLC is the maximal volume of gas within the lung, dividing it by the predicted lung weight provides an indication of maximal lung inflation. Total overinflated lung volume was calculated by summing the pixel area with a lung inflation value greater than the predicted maximal lung inflation value in each slice and multiplying by the slice thickness. This overinflation volume was expressed as a percentage of the total lung volume (ie, %overinflation).

The “upper lung zone” was defined as the region above the carina and zonal predominance was calculated using equation 5:

\[
\text{zonal predominance} = \frac{\text{upper overinflation (% of total upper lung)}}{\text{lower overinflation (% of total lower lung)}}
\]

The distribution was considered “upper zone predominant” if the result of equation 5 was \(>1\) and was considered “diffuse” or “lower zone predominant” if it was \(<1\). A cluster analysis was used to estimate the size distribution of the overinflated areas. The inverse slope of the log-log relationship of the size of the clusters (number of contiguous voxels that are inflated beyond the predicted value for maximum lung inflation for that individual) versus the number of clusters of that size is the power-law exponent (D). Individuals with diffuse small clusters of overinflated lung will have a steeper slope (ie, greater D) than individuals with larger overinflated regions. The low attenuation lung area (LAA) with an x ray attenuation lower than \(-950\) HU (%LAA\((-950))\) was calculated using the standard threshold approach and used to estimate “emphysema”. The zonal predominance and D were also calculated for %LAA\((-950))\).

Airway wall dimensions were measured for all visible airways cut in cross section on each CT image using the “full-width at half-maximum method”. Airway dimensions included lumen area (Ai), lumen perimeter (Pi), airway wall area (Aaw) and wall area expressed as the percentage of the total airway area (Aaw/A and Ai) = WA%) and a normalised airway wall estimate: square root of Aaw at Pi of 10 mm (ie, /Aaw at Pi10) (see online supplement). A mean (SD) of 33.2 (3.1) airways were measured per subject.

Statistical analysis

Statistical analyses were performed using JMP software Version 7.0.1 (SAS Institute, Cary, North Carolina, USA). The primary outcome was \(\Delta\text{FEV}_{1}\%\text{predicted/\text{year}}\) and the explanatory variables were CT measurements of (1) overinflated lung (ie, %overinflation, zonal predominance and cluster analysis); (2) emphysema (ie, %LAA\((-950))\), zonal predominance and cluster analysis); and (3) airway dimensions (ie, Ai, %WA and /Aaw at Pi10). Covariates included age, sex, body mass index (BMI), current smoking status (ie, current or ex-smoker), pack years and baseline spirometry measurements (FEV\(_1\)%predicted and FEV\(_1\)/FVC). Three multivariate models were used to identify the CT variables associated with the primary outcome after adjusting for the covariates (overinflation, emphysema and airway dimensions were tested respectively in models 1, 2 and 3).

To illustrate the relationship between \(\Delta\text{FEV}_{1}\%\text{predicted/\text{year}}\) and baseline %overinflation, we divided the 143 subjects into quartiles according to baseline %overinflation (quartile 1 had the least %overinflation) and compared \(\Delta\text{FEV}_{1}\%\text{predicted/\text{year}}\) across four quartiles using the Wilcoxon test. A linear mixed model was used to evaluate the annual decline in FEV\(_1\) (ml/year) for two groups (quartiles 1/2 and quartiles 3/4). Data were expressed as mean (SD) and p<0.05 was considered significant.
RESULTS

Baseline characteristics

Descriptive characteristics of 143 subjects are shown in Table 1 and baseline quantitative CT assessments are summarised in Table 2.

Follow-up measurements of FEV₁

Seventy-two of 143 subjects (50.3%) were seen twice over 2.3 (0.8) years, 49 (34.5%) were seen three times over 2.3 (1.1) years and 22 (15.4%) were seen more than three times over 3.5 (1.4) years. The mean (SD) number of follow-up visits was 2.7 (0.3) (range 2–5) over 2.5 (1.2) years (range 0.5–6.4). FEV₁%predicted/year observed over this time period averaged −2.0 (1.6)%/year (range −5.6 to +8.3)%/year.

Risk factors associated with annual change in FEV₁%predicted

Table 3 shows the three multivariate models testing the association between CT measurements and ΔFEV₁%predicted/year. In model 1, %overinflation was inversely associated with ΔFEV₁%predicted/year whereas neither emphysema nor airway dimensions were associated with ΔFEV₁%predicted/year in models 2 and 3. In addition, in model 1, sex and baseline spirometry measurements were also associated with ΔFEV₁%predicted/year (male sex: −0.73, 95% CI −1.43 to −0.03, p = 0.04; FEV₁%predicted: −0.16, 95% CI −0.11 to −0.22, p<0.01; FEV₁/FVC: 0.19, 95% CI 0.36 to 0.01, p = 0.03).

Overinflation and annual change in FEV₁%predicted

There was a significant linear relationship between CT %overinflation and ΔFEV₁%predicted/year (see online supplement). The mean (SD) baseline %overinflation for quartiles 1–4 were 37.1 (3.2), 52.6 (0.8), 67.0 (1.1) and 77.2 (0.6)%, respectively. The corresponding values of ΔFEV₁%predicted/year in the sex and baseline spirometry measurements were also associated with ΔFEV₁%predicted/year (male sex: −0.73, 95% CI −1.43 to −0.03, p = 0.04; FEV₁%predicted: −0.16, 95% CI −0.11 to −0.22, p<0.01; FEV₁/FVC: 0.19, 95% CI 0.36 to 0.01, p = 0.03).

DISCUSSION

The present results show a quantitative CT-based estimate of %overinflation using individual predicted maximal lung inflation is an independent predictor of rapid decline in lung function in smokers with normal baseline spirometry. The group with greater %overinflation at baseline exhibited a rate of decline in FEV₁ beyond the normal values predicted by Fletcher et al. These results suggest that, when the FEV₁ is normal, a quantitative structural assessment by CT can distinguish the smokers who will develop COPD from those who will not.

In the current study we used standard prediction equations for TLC and a prediction equation for lung weight to estimate maximal normal lung inflation for each individual. The mean (SD) value of predicted maximal normal inflation of the whole group was 5.9 (0.2) ml/g (range 5.3–6.5), of which the corresponding HU was −830 (5.5) (range −846 to −811), which is substantially different from the fixed cut-off value of 950 HU.

Table 2  Characteristics of baseline quantitative CT assessments of subjects

<table>
<thead>
<tr>
<th></th>
<th>Mean (SD)</th>
<th>Range</th>
</tr>
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<tbody>
<tr>
<td>Total lung volume(ml)</td>
<td>5086.2 (1350.3)</td>
<td>2600.1–9861.1</td>
</tr>
<tr>
<td>Mean lung density (g/ml)</td>
<td>0.2 (0.0)</td>
<td>0.1–0.2</td>
</tr>
<tr>
<td>Lung overinflation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>%overinflation</td>
<td>58.4 (16.4)</td>
<td>15.9–83.7</td>
</tr>
<tr>
<td>Cluster analysis</td>
<td>1.5 (0.2)</td>
<td>1.2–2.1</td>
</tr>
<tr>
<td>Upper zone predominant</td>
<td>n = 102</td>
<td></td>
</tr>
<tr>
<td>Diffuse distribution</td>
<td>n = 7</td>
<td></td>
</tr>
<tr>
<td>Lower zone predominant</td>
<td>n = 34</td>
<td></td>
</tr>
<tr>
<td>Emphysema</td>
<td></td>
<td></td>
</tr>
<tr>
<td>%LAA (−950 HU)</td>
<td>2.9 (2.6)</td>
<td>0.2–13.3</td>
</tr>
<tr>
<td>Cluster analysis</td>
<td>3.1 (0.4)</td>
<td>2.1–4.4</td>
</tr>
<tr>
<td>Upper zone predominant</td>
<td>n = 35</td>
<td></td>
</tr>
<tr>
<td>Diffuse distribution</td>
<td>n = 10</td>
<td></td>
</tr>
<tr>
<td>Lower zone predominant</td>
<td>n = 98</td>
<td></td>
</tr>
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</table>

Table 3  Multivariate models testing the association between the quantitative CT measurements and annual change in FEV₁%predicted

<table>
<thead>
<tr>
<th></th>
<th>Estimate (95% CI)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model 1*: Lung overinflation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>%overinflation</td>
<td>−0.04 (−0.08 to −0.02)</td>
<td>0.04</td>
</tr>
<tr>
<td>Cluster analysis</td>
<td>−0.49 (−11.3 to 10.35)</td>
<td>0.93</td>
</tr>
<tr>
<td>Upper zone predominant</td>
<td>0.02 (−0.10 to 0.14)</td>
<td>0.74</td>
</tr>
<tr>
<td>Model 2*: Emphysema</td>
<td></td>
<td></td>
</tr>
<tr>
<td>%LAA (−950 HU)</td>
<td>−0.13 (−0.47 to 0.21)</td>
<td>0.46</td>
</tr>
<tr>
<td>Cluster analysis</td>
<td>−0.61 (−2.68 to 1.45)</td>
<td>0.56</td>
</tr>
<tr>
<td>Upper zone predominant</td>
<td>0.11 (−1.38 to 1.59)</td>
<td>0.89</td>
</tr>
<tr>
<td>Model 3*: Airway dimensions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aaw (mm²)</td>
<td>0.17 (−0.40 to 0.73)</td>
<td>0.56</td>
</tr>
<tr>
<td>WA %</td>
<td>0.00 (−0.53 to 0.53)</td>
<td>0.99</td>
</tr>
<tr>
<td>Aaw at PI10 (mm)</td>
<td>4.4 (0.2)</td>
<td>3.9–5.1</td>
</tr>
</tbody>
</table>

*Models were adjusted for sex, age, body mass index, smoking status, pack years, FEV₁%predicted and FEV₁/FVC at baseline. Aaw, airway wall area; AI, lumen area; LAA, low attenuation lung area; Pi, lumen perimeter; WA, %, wall area expressed as the percentage of the total airway area. %overinflation, (volume of overinflated lung/volume of total lung) × 100%; %LAA (−950 HU), (volume of lung areas with x-ray attenuation lower than −950 HU/volume of total lung) × 100%; WA%, (airway wall area/airway wall area + lumen area)×100%; Aaw at PI10 (mm), Aaw for a standardised airway with an internal perimeter (ie, PI) of 10 mm.
2950 HU that is commonly used to define emphysema on CT scanning. Importantly, we do not claim that the minimally overinflated tissue identified by this procedure has undergone emphysematous destruction because we have no direct histological evidence. Based on pathology data, Leopold and Gough and McLean concluded that the dilation and destruction of the respiratory bronchioles that define centrilobular emphysema, the most common form of emphysema in smokers, is preceded by the disease in the terminal and preterminal bronchioles. We therefore strongly suspect that the minimal overinflation observed in this study may be caused by either minimal loss in the elastic recoil properties of the gas exchanging tissue and/or an increased resistance in terminal conducting airways due to inflammatory and tissue remodelling processes, both of which can occur before true emphysematous destruction.

The significance of the “overinflation” raised in our study is in agreement with that of the “hyperinflation” described by other investigators. Hyperinflation results from increased lung compliance due to emphysema and expiratory flow limitation, and patients may not perceive the negative results of it until an advanced stage because it develops slowly and insidiously over years. Ofir et al examined “hyperinflation” using lung function tests (total lung capacity, residual volume and functional residual volume) in patients with COPD GOLD stage I and healthy subjects. They found that patients with GOLD stage I COPD had more hyperinflation which increased as the intensity of dyspnoea increased. Casanova et al found that hyperinflation, independent of the BODE index, predicted mortality over a follow-up period of 34 months in 689 subjects with COPD. Although the findings that we have obtained with imaging tools are compatible with these studies of hyperinflation, further investigation is required to examine the relationship between these two measures of early disease.

Only a few studies have examined the relationship between quantitative CT measurements of the lung parenchyma and decline in lung function. Remy-Jardin et al examined 111 smokers and non-smokers and reported that subjects with emphysema visualised by radiologists at baseline had a more rapid decline in lung function than did those with normal CT scans. On the other hand, Parr et al and Stolk et al found no relationship between baseline CT emphysema and the subsequent decline in FEV1 in subjects with COPD. These studies recruited many subjects who already had moderate COPD at...
baseline, which contrasts sharply with the present study which was specifically designed to determine whether CT might identify those smokers who had normal initial spirometry and subsequently developed COPD.

In contrast to the extent of overinflation, the size and location of the overinflated regions, as assessed by cluster analysis and zonal predominance, was less helpful in identifying “susceptible smokers”. Moreover, although Nakano et al.22 showed that CT measurements of thickening and narrowing of the relatively large airways serve as a surrogate for the pathological changes in the small airways that are not measurable on CT scans, we failed to identify a relationship between the CT airway dimensions and ΔFEV1%predicted. There are several possible reasons for this observation. Most important, the differences in airway dimensions that accompany a relatively small change in lung function are probably beyond the resolution of CT scans. Although many investigators have shown relationships between airway wall dimensions and airflow obstruction in cross-sectional studies, the range of lung function in those studies was much larger than in the present study.23 Second, quantitative histological studies have shown that statistically significant airway wall thickening does not become apparent until the later stages of disease (GOLD stages III and IV) with an FEV1%predicted <50.24 Finally, our method for measuring airway dimensions may not be optimal for assessing subtle changes. Hasegawa et al.25 used volumetric scanning to show that airflow limitation in COPD is more closely related to the dimensions of the distal smaller airways (ie, 5th and 6th generations) than those of proximal larger airways (ie, 3rd and 4th generations).

Fletcher et al.26 were the first to show a relationship between the initial FEV1 and its subsequent decline and referred to the phenomenon as the “horse racing” effect. However, Burrows et al.27 observed an opposite relationship between ΔFEV1 and initial FEV1, such that the higher the initial FEV1, the more negative the ΔFEV1. They pointed out that this was due to “regression toward the mean”, a phenomenon in which subjects performing especially well on their first test show a greater decline because of a poorer performance on subsequent tests. Burrows and Stanescu27 also observed an association between initial FEV1/FVC and ΔFEV1%predicted, and suggested that FEV1/FVC might provide a more reliable indicator of future loss in FEV1. Our results confirm the findings of Burrows and Stanescu and extend their observations by showing that CT evidence of lung parenchymal overinflation is an independent predictor of decline in FEV1.

Although the lack of an association between smoking intensity (ie, pack years) and decline in lung function might be a little surprising, this is consistent with other reports in the literature.28 We also think this lack of association between pack years and the decline in FEV1 supports Fletcher’s concept that “non-susceptible smokers” remain unaffected regardless of their smoking history. The converse is also true—that “susceptible smokers” exhibit a decline in lung function independent of their smoking status.

A limitation of this study is that it was not originally designed as a study of COPD but was added on to a lung cancer screening cohort. Subjects were therefore not randomly chosen from the population. Follow-up spirometry was arranged at the time of follow-up CT scans which depended on the characteristics of the lung nodule(s) found on the initial CT scan. This means that different numbers of spirometric tests were performed at different frequencies between subjects. To overcome this limitation we used ΔFEV1%predicted/year also adjusted for the normal annual loss due to ageing and corrected for differences between women and men.

In summary, we conclude that the quantitative assessment of the lung inflated beyond individually predicted maximal lung inflation on initial CT scans may be able to identify the “susceptible minority of smokers” who eventually will develop COPD. Our working hypothesis is that the minimally overinflated lung contains the earliest forms of lesions that either increase peripheral airway resistance and/or increase lung compliance by initiating emphysematous destruction.

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Authors’ contributions: All authors contributed to the design of the study and the drafting and reviewing of the manuscript. RY, YN and JCW performed the laboratory work and statistical analysis. JCH, PDP, HDC, DDS and SL are the principal investigators of the project, obtained funding for and supervised the project. SL and AMMcW initially recruited the subjects and DDS provided statistical suggestions. All authors read and approved the final manuscript.

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Competing interests: JCH has served as a consultant, given lectures and participated in advisory boards of several major pharmaceutical companies in the past five years. The total reimbursement for these activities is less than $20,000. PDP was the principal investigator of a Merck Frosst supported research programme to investigate gene expression in the lungs of patients who have COPD. He and collaborators have received approximately $200,000 for this project. These funds have supported the technical personnel and expenses involved in the project. He sits on an advisory board for Tacleis Biotherapeutics who make anti-one antitrypsin replacement therapy. He is the principal investigator of a project funded by GlaxoSmithKline to develop CT-based algorithms to quantify emphysema and airway disease in COPD. With collaborators he has received approximately $300,000 to develop and validate these techniques. These funds have been applied solely to the research to support programmers and technicians.

DOS has received research funding from GlaxoSmithKline and AstraZeneca for projects on chronic obstruction pulmonary disease. He has also received honoraria for speaking engagements for talks on COPD sponsored by these organizations. HDC received $11,000 in 2005 and $4800 in 2006 and 2007 for serving on an advisory board for GlaxoSmithKline. He is co-investigator on two multicentre studies sponsored by GlaxoSmithKline and has received travel expenses to attend meetings related to the project. He has three contract service agreements with GlaxoSmithKline to quantify the CT scans in subjects with COPD and a service agreement with Spirion Inc to measure changes in lung volume in subjects with severe emphysema. A percentage of HDC’s salary between 2003 and 2006 (15 000 US $/year) derives from contract funds provided to a colleague PDP by GlaxoSmithKline for the development of validated methods to measure emphysema and airway disease using computed tomography. HDC is the co-investigator (with DDS) on a Canadian Institutes of Health-Industry (Wyeth) partnership grant. There is no financial relationship between any industry and the current study. RY, JCW, YN, SL and AMMcW have no competing interests in the content of this paper.

Ethics approval: The University of British Columbia Clinical Ethics Review board approved the study and all subjects provided informed written consent for the use of all materials and data.

Provenance and peer review: Not commissioned; externally peer reviewed.

REFERENCES

The sst1 locus controls granuloma necrosis in tuberculosis

There is broad variation in susceptibility to *Mycobacterium tuberculosis* (MTB), and mouse models play a key role in elucidating the underlying mechanisms. The authors tested the hypothesis that caseation within pulmonary lesions is a specific effect of the sst1 (supersusceptibility to tuberculosis 1) locus on chromosome 1.

Typical mouse lesions described in the literature lack central caseation. However, the C3HeB/FeJ strain of mouse develops large necrotising granulomas after exposure to MTB. The authors compared the course of infection in an MTB-resistant mouse strain (B6) with an sst1-susceptible congenic strain (B6.C3H-sst1) which was genetically identical except for the interval containing the sst1 locus.

They found that, although initial dissemination was similar, bacterial loads, clinical disease, lung necrosis and mortality were considerably worse in the strain containing the sst1-susceptible allele. In addition, relapse after 8 months of isoniazid was faster and more rampant in this group. After demonstrating significantly slower disease progression and milder histopathology in the B6.C3H-sst1 strain compared with the C3HeB/FeJ strain, the authors concluded that the effect of the sst1 locus is modified by the genetic background of the host. Enhanced pro-inflammatory cytokine production by macrophages was observed in the susceptible strains; however, this appears to be controlled by loci other than sst1.

Further characterisation of sst1-encoded molecular mechanisms may not only shed light on a key aspect of the pathogenesis of tuberculosis, but may also suggest therapeutic interventions to reduce lung pathology and transmission of the pathogen.


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