Significant intimal abnormalities in muscular pulmonary arteries of patients with early obstructive lung disease

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SUMMARY Substantial intimal changes, unrelated to aging, were found in resected lobes or lungs of 30 smokers with early obstructive lung disease (22 men, eight women; age range 46–74 years; forced expiratory volume in one second (FEV1) % predicted range 47–119%). Intimal area was measured using a digitiser and expressed as a proportion of the area enclosed by the internal elastic lamina (IEL), correcting for constriction (intima index). Arteries were subdivided into four size (length IEL) groups. For the smallest group (≤600 μm length IEL) the mean intima index (II600) ranged from 0.09–0.34; half the values were ≥0.20. Intimal abnormality was unrelated to the size or site of tumour or to macroscopic emphysema, and correlated weakly with pack years. II600 values were, however, significantly correlated with factors known to be affected by smoking such as alveolar wall surface area per unit lung volume, FEV1 % predicted, and FEV1/FVC (forced vital capacity); these last two factors were also significantly associated with intimal abnormality in arteries measuring 601–1200 μm length IEL.

An increase in the intima of muscular pulmonary arteries may have functional implications for the pulmonary circulation in terms of flow and pressure changes.1–3 As such there have been many studies of the intima both in normal and disease states. Those studying healthy subjects have commented that patchy intimal changes are a common and very variable feature of aging.4–7 The issue of whether smoking has an effect on the intima is still open to debate. Some workers have implicated smoking as a cause of increased intimal thickening in muscular pulmonary arteries.8–10 In a study of 23 subjects without evidence of any cardiopulmonary disease likely to have affected the pulmonary vasculature, however, we could find no systematic differences in the intima between smokers and non-smokers.9 Although the extent of intimal abnormality is very variable in healthy subjects, it is generally acknowledged that clinically important widespread abnormality is normally found only in association with specific pathology.13

This paper describes the pronounced intimal abnormality observed in the resected lobes and lungs of 30 smokers with small peripheral lung carcinomas, and how this relates to structural and functional abnormalities within the lung.

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Material and methods

Between September 1980 and May 1984 material was obtained from resection operations carried out at the City Hospital, Edinburgh, on patients with suspected lung carcinomas. Thirty of these specimens were included in the present study; three were whole lungs and 27 were single lobes (18 upper, three middle, and six lower lobes).

The patients were predominantly male (22 men, eight women ranging in age from 46 to 74 years; all were current (n = 26) or ex-smokers (n = 4). Data were available on amount smoked, years smoked, and years of not smoking (where relevant) and were used to calculate total pack years. In all cases the smoking histories were obtained directly from the patient using a standard questionnaire.

During the week before surgery, patients underwent a series of pulmonary function tests and measurements of blood gases. The data available included: forced expiratory volume in one second (FEV1), forced vital capacity (FVC), total lung volume (TLC), residual volume (RV), transfer factor (TCO), transfer factor per unit of lung volume (KCO), PaO2 and PaCO2. The FEV1 and FVC measurements were additionally expressed as a percentage of predicted values for age and height for women14 and for men.11
**Significant intimal abnormalities in lungs of smokers**

**Preparation of Specimens**
After inflation with formal saline through the main or lobar bronchus fixation was allowed to continue for a minimum of 24 hours before cutting lungs or lobes into 1 cm sagittal slices. Twelve tissue blocks (1·9 × 1·9 × 0·6 cm) were then taken from the two most lateral slices. Areas containing carcinomas, which were generally small and peripherally situated, were avoided. The 12 blocks, which were selected from the remainder of these two slices, were selected using a random sampling technique. After embedding in glycol methacrylate the tissue blocks were sectioned at 3 μm using the technique of Sims and stained for elastic.

**Method of Measurement**
Cross sectionally cut muscular pulmonary arteries were assessed by measuring the area of the intima and relating this to artery size using total length of internal elastic lamina (IEL) as the criterion for artery size. All measurements were obtained using a semiautomatic digitising system; precise details of the measuring procedures followed have been reported previously. An average of 57 muscular pulmonary arteries were measured in each subject.

**Calculation of Intima Indices**
Data on the intima of each measured artery were expressed in a way which reflected the true extent of lumen occlusion (the intima index). These intima indices were calculated by dividing the area of the intima by the area enclosed by the IEL in its theoretically uncollapsed/unconstricted state. This is necessary to accommodate differences in the degree of collapse or constriction between arteries, which can be very variable. Values for intima index range from >0 to ≤1, indicating minimal through to total occlusion of the artery lumen by intimal change.

**Assessment of Emphysema**
Emphysema was assessed macroscopically as a percentage of the area of the mid-sagittal lung or lobe slice. Types of emphysema were not considered separately.

Emphysema was also assessed microscopically by a method which assesses alveolar wall surface area per unit volume (AWUV as mm²/mm³) from measurements of alveolar perimeter per unit area on histological sections using an IBAS2 automatic analyser. Measurements were made on random fields (18–35) from each of the 12 tissue sections available from each subject. Perimeter measurements were converted to AWUV by the formula AWUV = alveolar perimeter (mm/mm²) × 4 ÷ π. In our study the AWUV value used to represent each subject was that of mean AWUV.

**Analysis of Data**
The data on individual arteries together with those on smoking, respiratory function, blood gases and emphysema were transferred to tape on the mainframe computer (a PRIME 750) at the Institute of Occupational Medicine. All analyses were carried out using the statistical package "Minitab".

**Results**

**Preliminary Analysis of Intima Indices for Individual Subjects**
For all subjects the intima index was found to vary with artery size (total length of IEL), smaller arteries being proportionately more affected by intimal change (fig 1). To compare subjects it was therefore necessary to calculate mean intima indices for arteries subdivided into size groups. In previous studies the size groups chosen were: up to 600 μm total length of IEL, 601–1200, 1201–1800, and >1800 μm total length of IEL. Analysis of the relation between intima index and artery size for all 30 subjects showed that these four size groups were all the most appropriate to use for the present study. The corresponding mean intima indices were termed II600, II1200, II1800 and II >1800, the figures denoting the upper limit of the size range of arteries included in each group.

The range of the mean intima indices observed in the 30 resection specimens was as follows: II600 0·09–0·34; II1200 0·06–0·24; II1800 0·03–0·13; II >1800 0·02–0·07.

**Relation Between Intimal Abnormality and the Study Variables**
In the resection specimens intimal abnormality was not related to age—shown for the smallest size group of arteries (II600) in fig 2 (r = 0·14). The value for the II600 ranged from 0·09–0·34, and more than half of

![](http://jcp.bmj.com/)

**Fig 1** Relation between intima index and artery size for one subject.
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Fig 2  Relation between mean intima index of arteries in smallest size group (II600) and age.

the patients had values of 0·2 or greater—that is, greater than 20% lumen occlusion.

Although positive associations were observed between smoking (expressed as pack years) and intimal abnormality (expressed as mean intima indices), the associations were heavily influenced by a very small number of subjects with high values for pack years. The II600 result is illustrated in fig 3.

In terms of structural changes within the lung there was no association between intimal abnormality and macroscopic emphysema, the incidence of which in the study group was low. A significant (p < 0·02) positive association, however, was observed between intimal abnormality and mean AWUV, specifically for the smallest size group of arteries (fig 4). A low AWUV value indicates a loss of alveolar surface area—that is, microscopic emphysema. In the resection specimens low AWUVs tended to be found in association with high levels of intimal abnormality and vice versa.

Intimal abnormality was also significantly associated with functional as well as structural changes within the lung. Values for II600 (fig 5) and III1200 were both significantly (p < 0·02) correlated with FEV1, expressed as a percentage of that predicted for age and height. Similar significant associations were observed between intimal abnormality (II600 and III1200) and FEV1, expressed as a percentage of FVC (results not shown). The r values were −0·61 and −0·48, respectively.

Intimal abnormality was unrelated to any of the other pulmonary function or blood gases data.

Discussion

This study has shown that there is a significant level of intimal abnormality in a group of smokers, which is associated with some degree of structural and functional abnormality within the lung.

It is appropriate perhaps to begin a discussion of these findings with some comments on the material used in the study. Firstly, our material comprised a mixture of whole lungs and lobes and as such we were not comparing subjects on a strictly similar basis, but we felt that this was justifiable on the grounds that we have previously shown that intimal abnormality does not vary between lobes.9 Secondly, the subjects studied were a highly selected group in that they all had peripheral carcinomas and were considered to be fit for operation. This leads to one possible explanation for the pronounced intimal abnormality: it is somehow related to the presence of the carcinoma. Tissue blocks taken distal to the segment containing the carcinoma, however, showed no evidence of more severe intimal abnormality, and from this we concluded that the tumours were not having a local effect on the intima.

Other workers have studied pulmonary blood vessels in resection specimens.20 21 Some concluded that the increased arterial intimal thickening was associated with smoking.21 The other group, who specifically studied the ultrastructure of the intimal fibrosis, believed the intimal fibrosis to be associated with aging rather than the bronchial neoplasm.20 Our results indicate that the intimal abnormality is unlikely to be simply a function of aging. First there was no direct relation between the level of intimal abnormality and patient age, which may be due to the fact that the study group was predominantly middle-aged and did not cover the extremes of age. A second factor which militates against aging being solely responsible is that the extent of intimal abnormality in the resection specimens was much higher than that observed in a previously studied group in whom we detected an age effect.9 That particular necropsy study

Fig 3  Relation between mean intima index of arteries in smallest size group (II600) and pack years.

Fig 4  Relation between mean intima index of arteries in smallest size group (II600) and microscopic emphysema (AWUV).
Fig 5  Relation between mean intima index of arteries in smallest size group (II600) and FEV₁, expressed as percentage of predicted value.

consisted of subjects showing no evidence of any cardiopulmonary disease likely to have affected the pulmonary vasculature. The differences between these two study groups are considerable and are quite specific to the intima. Taking the smallest size group of arteries as an example, only one of the 23 subjects in the group without any cardiopulmonary disease had an II600 value of \( > 0.2^\circ \) whereas more than half of the 30 subjects in the resection group did so.

We consider lumen occlusion in excess of 20% to be quite pronounced given that these values are minimum possible values because they were calculated independently of artery constriction or collapse to ensure comparability between subjects. If the degree of lumen occlusion seen in the internal elastic lamina does reflect the amount of tone in the arteries during life, and is not simply an artefact after death, then the degree of lumen occlusion by intimal abnormality is very much higher. To illustrate this, one of the patients in the resection group had a mean II600 of 0.32 after accounting for constriction or collapse—that is 32% lumen reduction in arteries measuring less than 600 \( \mu \)m length of internal elastic lamina. If the constriction collapse had not been accounted for, the observed actual mean II600 would have been 0.72 or 72% lumen reduction. Such severe lumen reduction has obvious implications with respect to pulmonary vascular resistance.

The differences we have noted in intimal abnormality between the group without any cardiopulmonary disease and the resection group are probably related to smoking, either directly or indirectly, those in the resection group being, in general, much heavier smokers. We have shown that the level of intimal abnormality, particularly in the smaller arteries, is strongly correlated with both structural (reduced alveolar surface area per unit lung volume) and functional (reduced ventilatory capacity) abnormalities within the lung, both of which are known to be related to smoking. It may be that the resection group comprises patients with a particularly high response to smoking.

In general, the patients we studied had evidence of early obstructive lung disease. Given the nature of the study group, however, some of the patients will probably go on to develop more severe obstructive lung disease. Our results are interesting in that they indicated that pronounced vascular (intimal) changes occur relatively early in the natural history of chronic airflow obstruction. Although the precise functional importance of these vascular changes is not known, they may be important in the eventual development of pulmonary hypertension.

References

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