Vertical gradients of lung density in supine subjects with fibrosing alveolitis or pulmonary emphysema

A B Millar, D M Denison

Abstract
Computed tomography was used to determine the vertical gradient of density in the peripheral lung tissue of 12 patients with histologically proved fibrosing alveolitis and 12 patients with chronic bronchitis and evidence of pulmonary emphysema on the computed tomodograms. Measurements were made at total lung capacity and at residual volume and compared with similar measurements from 12 normal subjects reported in a previous study. At residual volume the mean peripheral tissue density in the emphysematous lungs was 0.081 kg/l compared with 0.426 kg/l in the fibrotic lungs and 0.323 kg/l in the normal lungs. The observed densities in the three groups were compared with those in a theoretical model predicting the vertical changes of lung density caused by gravitational effects that would be found in lungs with differing compliance. The emphysematous lungs showed a much greater increase of density with descent down the lung than that predicted for normal lungs, and the results were explicable by an increase in compliance. The fibrotic lungs showed considerably less change in density than expected, implying loss of compliance. It is suggested that local changes of compliance are important determinants of vertical density gradients in diseased lungs.

We reported previously that measurements of area and density in thoracic computed tomodograms taken at total lung capacity (TLC) and residual volume (RV) can be used to assess regional gas and tissue volumes and ventilation in normal and diseased lungs.1,2 We have also investigated the vertical density gradients in the lungs of normal subjects and suggested that they were related to changes in local compliance.3 This interpretation was based on the model of the lung in which this behaves as a liquid, open cell foam collapsing under its own weight, as proposed by Wilson and Bachofen in 19824 and confirmed recently by Stamenovic and Yager.5 Other authors have already commented on the alterations in the vertical density gradients present in thoracic computed tomodograms of patients with emphysema6-8 or fibrotic lung disease,9 as illustrated in figure 1. We have studied such changes in more detail using information present in the scans and in the lung function results of 12 patients with cryptogenic fibrosing alveolitis and 12 patients who met the Medical Research Council criteria for chronic bronchitis10 and who also had evidence of emphysema on their scans. Other aspects of their scans have been reported previously.2

Methods

SUBJECTS AND LUNG MEASUREMENTS
The subjects were selected retrospectively from large groups of patients considered to have probable emphysema or fibrosing alveolitis who had computed tomography performed for clinical reasons. Each patient had detailed lung function tests within 24 hours of their scan. The methods used and normal values have been reported previously.1,2 The 12 patients with fibrosing alveolitis had histologically proved disease. There were eight men and four women, their ages ranging from 21 to 69 years and their height from 161 to 183 cm. Their lung function, expressed as mean (SD) % predicted, was as follows: forced expiratory volume in one second (FEV1) 66 (17); forced vital capacity (FVC) 65 (11); residual volume (RV) 58 (13); single breath gas transfer factor (TLCO) 35 (9); transfer coefficient (KCO) 66 (18). The other 12 patients (two women and 10 men; age 43–75 years, height 163–187 cm) met the Medical Research Council criteria for chronic bronchitis10 and had computed tomography evidence of widespread emphysema. Their lung function test results were as follows: mean (SD) % predicted: FEV1, 38 (23); FVC 77 (26); TLC 133 (13); RV 220 (63); TLCO 43 (18); KCO 55 (18).

All the computed tomodgrams were obtained as the patient lay supine. We used an Elscint 2002 whole body scanner, which has a single scan time of five seconds. Data from the scanner were stored on magnetic tape and transferred to a separate viewing console for analysis. The protocol for scanning each subject was identical to that previously described.3 The subject was asked to breathe in to TLC and signal when the lungs were full. An inspiratory scan was recorded. This procedure was repeated from lung apex to lung base, with a constant slice width of 1 cm. At the end of inspiratory scans corresponding information was obtained for the lungs at residual volume, the patient being asked to breathe out and signal when the lungs were “empty.” The mean densities of successive horizontal planes of peripheral lung tissue were calculated by

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Figure 1 Computed tomogram of a supine subject with emphysema (a) or fibrosing alveolitis (b) at full inspiration (TLC) in the upper scans and full expiration (RV) in the lower scans, showing the differences in vertical density gradient.

summing values from the equivalent levels in each scan, the mid point of the right atrium being used as the reference level. The mean (SD) densities of each 1 cm thick plane of peripheral lung tissue are presented. The densities observed at residual volume in the two groups of patients are compared in figure 2 with those previously reported for normal subjects.

ANALYSIS

We described previously a simple mathematical model of a lung like foam to predict the vertical density gradient of normal lung. In that model the compliance of the lung was assumed to be constant from the top to the bottom of a stack of lung tissue for a given degree of lung inflation. We wished to improve on that model by allowing for variations in lung compliance due to different degrees of inflation within any stack. To do this we began graphically, by differentiating the pressure-volume curve of normal lung—for example, that published by Murray—to obtain the compliance for each degree of inflation from RV to TLC. Normalised for lung volume, the differential was best fitted by the curve shown in figure 3a, for which the equation is:

\[ CL = 5.7 - (2 \times \frac{VL}{TLC}) - (\frac{VL}{TLC})^{2} \times \text{TLC/cm H}_2\text{O}. \]

A synthetic pressure-volume relationship for normal lung obtained by integrating this equation is shown in figure 3b. Armed with this equation we calculated the vertical distribution of density in a tall stack of lung tissue collapsing under its own weight. The upper element of the stack was taken to be at
101% of TLC. We assumed the lung characteristics described by Weibel, with a weight of 825 g and a TLC of 6400 ml. At any given degree of whole lung inflation the vertical density gradient within the lung would be represented by some segment of this curve.

Previously we noted that the supine lung is about 17 cm tall at TLC and shrinks to some 12 cm at RV. When the stack of lung tissue is related to the mid point of the right atrium, 1 cm of this reduction in height comes from the top of the stack and 4 cm from the bottom. Using that information allows us to take appropriately sized segments from the synthetic curve described above and plot them as the “family” of vertical density gradients to be expected in computed tomograms of normal lungs at various degrees of inflation from TLC to less than normal RV. A selection of such predicted normal vertical density gradients at various degrees of inflation (from greater than TLC on the left to less than RV on the extreme right) is shown by the solid lines in figure 4. We have superimposed the real data of the patients at RV from figure 2 and data from the 12 normal subjects from reference 3.

Results

The mean peripheral lung density in the emphysematous lungs was 0.061 kg/l at TLC. This changed little with reduction in lung volume and was 0.081 kg/l at RV (table 1). The density values ranged from 0.039 kg/l to 0.103 kg/l at TLC in the 12 subjects and from 0.048 kg/l to 0.142 kg/l at RV. The gradient was greater than would be predicted by comparison with the “family” of calculated density gradients shown by the solid lines in figure 4.

In the lungs of patients with fibrosing alveolitis the mean peripheral lung density was 0.211 kg/l at TLC and 0.426 kg/l at RV. The range of density values in the 12 subjects was from 0.151 to 0.253 kg/l at TLC and from 0.295 to 0.620 kg/l at RV. The gradient was less than would be predicted by comparison with the family of curves shown in figure 4.

Discussion

We have shown previously that in normally compliant lung the greater the density the more pronounced the curve—that is, the density gradient. Density in normal lung is directly related to the degree of inflation; thus a vertical density gradient predicted for normal lung at TLC would be to the left and at RV to the right of the curves for normal lung shown in figure 4. Although the patients with emphysema had lightweight lungs and a low KCO, suggesting reduced pulmonary blood volume, their lungs show a much greater than predicted increase in density towards the bottom of the supine lung, reflecting a substantial increase in compliance. The patients with fibrosis on the other hand, with quite similar losses of pulmonary capillary blood volume, show a less than predicted increase in density with descent down the lung. Their observed gradient is similar to that in the normal subjects with a lower mean density, indicating a considerable loss of compliance.

Comparisons of computed tomograms with lung models and with isolated healthy and diseased lungs show that tomograms provide accurate maps of physical density distributions. Many authors, such as Kree, observing the vertical density gradients in healthy lungs, have noted that they are more obvious in expiration than inspiration (when most lung units are fully inflated on the plateau of the pressure-volume curve and hence have similar compliance) and that they are inverted with reversal of posture, and have attributed them to the effects of gravity on the pulmonary circulation. Alterations of the vertical density gradients in diseased lungs have also been observed frequently, alterations again being principally attributed to loss, distension, or leakage of vessels. We suggest that vertical density gradients also reflect local lung compliance, especially in expiration.

Global lung compliance was measured in
our laboratory in 15 normal subjects, 15 patients with emphysema, and 15 patients with fibrosing alveolitis, the three groups yielding mean values of 0.23, 0.74, and 0.12 1/cm H₂O (Ward et al, unpublished observations). These values compare well with those published elsewhere.16-20 Computed tomography confirms that emphysema and fibrosis are not uniformly distributed throughout the lung. Consequently, measurements of whole lung compliance substantially underestimate the severity of changes in the damaged parts of the lung. If, for example, the 15 patients with emphysema mentioned earlier had half of their lungs affected the compliance of the emphysematous half would be 1.25 1/cm H₂O—that is, a fivefold increase. Similarly, if the 15 patients with fibrosing alveolitis had half of their lungs affected the compliance of the fibrotic part would be 0.01 1/cm H₂O, a more than 10 fold drop. We believe that it is these severe local changes as well as the effects of gravity on blood vessels that contribute to vertical density gradients in computed tomograms of diseased lungs.

Figure 4  Mean densities observed at residual volume in 12 patients with emphysema (squares), 12 patients with fibrosing alveolitis (triangles), and 12 normal subjects (circles; from a previous study) compared with predicted densities described in the text (solid lines).

<table>
<thead>
<tr>
<th>Mean (SD) density (kg/l)</th>
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<tbody>
<tr>
<td>Height (cm) above (+) or below (−) mid right atrium</td>
</tr>
<tr>
<td>Right</td>
</tr>
<tr>
<td>+5</td>
</tr>
<tr>
<td>+3</td>
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<td>+1</td>
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<td>−7</td>
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<td>−10</td>
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<td>−12</td>
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Table 1  Density of cephalocaudal samples of peripheral lung tissue in 12 supine subjects with emphysema

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