Lung mechanics in subjects showing increased residual volume without bronchial obstruction

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ABSTRACT Fourteen subjects showing an increase of residual volume (RV) without any clinical or functional signs of bronchial obstruction were studied. Maximum expiratory flow volume (MEFV) curves were obtained with a pressure-corrected volume plethysmograph. Static pressure-volume curves were obtained by stepwise interruption of a slow expiration from total lung capacity (TLC) to RV. Static compliance was measured by the slope of pressure-volume curve between functional residual capacity (FRC) and FRC+20% of TLC. Maximum flow static recoil (MFSR) curves were constructed by plotting MEF obtained from MEFV curves against elastic pressure (Pst) obtained from pressure-volume curves at the same lung volumes. Most patients demonstrated a decrease of MEF 50% and 25% of VC. From the MFSR curves it was clear that this reduction was not the result of increased airways resistance, but rather of loss of elastic recoil. Most patients showed a significant decrease of Pst at different volumes and changes seem likely to be evidence of emphysema.

Residual volume is determined by the point at which the pressure produced by the muscles of expiration is dissipated entirely on the elastic recoil of the respiratory system. Increase in residual volume may result from loss of elastic recoil, any form of airways obstruction, or both. The purpose of this work was to study some aspects of lung mechanics in subjects with an increased residual volume in the absence of any evidence of bronchial obstruction.

Methods

We examined 14 patients, 13 men and one woman, with a mean age of 45.5±8.2 years (range 29–56 yr). These subjects complained of minor respiratory disability, mainly slight exertional dyspnoea. All save one were smokers—three smoked less than 10 cigarettes per day, the remainder 15–20.

Clinical examination showed no signs of cardiac disease or bronchial obstruction and all chest radiographs were considered to be normal. Study of respiratory function showed only an increase, in comparison with normal values,1 of residual volume measured by plethysmography, the average increase being 58.14±26.33 (range: +32–104%).

Vital capacity (VC), forced expiratory volume in one second (FEV1), forced expiratory volume in one second/vital capacity ratio ×100 (FEV1/VC), and airways resistance (Raw) were close to predicted values in every patient (table 1).

Vital capacity and FEV1 were measured with a pneumotachograph (Jaeger) coupled to a pressure transducer (Jaeger) and were obtained by integration of the flow measured at the mouth and recorded on an X-Y recorder. The result was taken as the best of two or three efforts and the volumes, at BTPS, were expressed as a percentage of the predicted normal values for sex, height, and age.2 Functional residual capacity was determined by a constant volume body plethysmograph (Pneumotest-Bodystat Jaeger, Würzburg) which also allowed, with a single manoeuvre, the simultaneous measurement of VC. Although the spirometric and plethysmographic values of VC were almost identical we calculated RV from plethysmography. Functional residual capacity was measured in duplicate. An approximation of 250 ml or less was required for two successive determinations. From the FRC, TLC, RV, and RV/TLC were calculated.

We adopted the normal values of Amrein et al1 for RV, FRC, TLC, and RV/TLC ratio. These values are not significantly different from those

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of Cotes,\textsuperscript{2} except for RV/TLc which is significantly lower (p<0.05). In our 14 cases the predicted mean values of RV/TLc was 27±3·10 according to Amrein \textit{et al} and 31±2·8 by Cotes. The functional residual capacity and RV of every patient were expressed as a percentage of the respective predicted values.

Airways resistance (Raw) was obtained during tidal breathing by the slope of the line traced between inspiratory and expiratory flows of 0.5 l s\(^{-1}\). Five measures were made in sequence and we calculated the average of the last three. The predicted values of Raw are those of Amrein \textit{et al}.\textsuperscript{1}

Maximum expiratory flow volume (MEFV) curves were obtained with the patient sitting in the pressure-corrected, volume plethysmograph (Pneumotest-Bodytest Jaeger, Würzburg). To relate maximum flow rates to lung static recoil we chose the plethysmographic volume measurements, since static recoil pressure depends on the actual lung volumes, regardless of alveolar gas compression.\textsuperscript{3} An appropriate device prevents the response time of the flow channel exceeding that of the volume recording channel. Volume change measured in the plethysmograph was recorded on the abscissa, and flow, measured at the mouth, on the ordinate. The curves were registered on a X-Y recorder. The best of three or four attempts was chosen for the measurement of MEF. The curve was quantified by measuring MEF at lung inflation of 75%, 50%, and 25% of VC. As normal reference values we adopted those of Cherniack \textit{et al}.\textsuperscript{4}

Intraoesophageal pressure was measured using the method of Milic-Emili \textit{et al}.\textsuperscript{6} An oesophageal balloon (length 9 cm, wall thickness about 0.04 mm) was placed in the middle third of the oesophagus and tested for freedom from artefacts. With the patient sitting in the plethysmograph, three or four static pressure–volume curves were obtained by stepwise interruption of a slow expiration from TLC to RV. Transpulmonary pressure (Pst), expressed as the difference between mouth and oesophageal pressure, was measured with a differential pressure transducer (Jaeger). These curves were registered on an X-Y recorder, producing volume change on the ordinate and Pst on the abscissa. The "best" of them was used to measure Pst at different volumes. The "best curve" covers a range of volume, from TLC to RV, as close as possible to VC of the patient and is free of potential errors in elastic recoil estimation, which are mainly caused by active oesophaegal contractions, arising while or shortly before recording the curve.

Values of Pst at different volumes and of static compliance (Cstat) were found to be reproducible. Static pressure–volume curves were obtained by the above-mentioned method on five different days in eight normal subjects, aged from 20–24 years. When the above criteria were not satisfied, the curves were discarded and the study was repeated on the following days. A total of 240 valid measurements was, therefore, obtained in these subjects, the six values Cstat, Pst 100%, 90%, 80%, 70%, and 60% of TLC for each of the five days. Mean values of the five measures of Cstat and Pst at different volumes are given in table 2. In every subject all values for the same measurement fell within the range of ±2 SD.

Predicted values for Pst were derived from these data of Yernault \textit{et al}.\textsuperscript{8} Static compliance was measured by the slope of the pressure–volume curve between FRC and FRC+20% of TLC.
Predicted values of Cstat, as a function of height, are those proposed by Yernault and Engert.\(^7\)

Maximum flow static recoil (MFSR) curves\(^8\) were constructed by plotting MEFV obtained from MEFV curves against corresponding PsT calculated from static pulmonary pressure–volume curves at the same lung volumes.

Table 3 shows the maximum expiratory flow at 75%, 50%, and 25% of vital capacity. Mean values and standard deviation in 14 subjects showing only increased residual volume in the absence of signs of bronchial obstruction.

The MEFV values of 14 patients, calculated from the flow volume loop at 75%, 50%, and 25% of VC, are given in table 3. Most patients had decreased MEFV values at different volumes. In comparison with the predicted values, the average decrease of MEFV 50% and MEFV 25% was highly significant (p<0.001). On the other hand the decrease of MEFV 75% was not significant.

It should be emphasised that, unlike MEFV, for which predicted values are usually given as a percentage of VC, normal reference data of PsT, as a rule, are indicated as a percentage of TLC, and we have followed this convention in the statistical evaluation of our results.

Static compliance and PsT values at 100%, 90%, 80%, 70%, and 60% of TLC are given in table 4. Ten of 14 subjects showed an increase of Cstat, but there was no significant difference between the mean value of Cstat obtained in these individuals and the mean predicted value. Most patients however showed a decrease of PsT at different volumes of TLC. As a rule, the mean value of PsT measured at a given volume was significantly decreased in comparison with the mean predicted normal value, with the exception of PsT 100% of TLC.

The figure shows the maximum flow-static recoil (MFSR) curves of the 14 patients examined. Since the curves are constructed by plotting MEF against PsT at the same lung volumes, both of them are in this case expressed as a percentage of VC. Because MEF varies with lung size, flow in 1 sec\(^{-1}\) at different volumes was divided in each patient, according to Mead et al.\(^8\) by individual TLC and plotted on the ordinate. Transpulmonary pressure, expressed as a percentage of VC (table 5), is plotted on the abscissa.

The figure also presents three MFSR curves, derived from Mead et al.\(^8\) each being the mean curve of five normal subjects in the age groups 24–30, 38–43, and 48–61 yr, and, as an example,
Table 4  Static compliance (Cstat) and transpulmonary pressure (Pst) at 100%, 90%, 80%, 70%, 60% of TLC* and mean values ± SD in 14 subjects showing only increased residual volume

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<th>Subjects</th>
<th>Cstat (kPa)</th>
<th>Pst 100% TLC (kPa)</th>
<th>Observed/Predicted†</th>
<th>Pst 90% TLC (kPa)</th>
<th>Observed/Predicted†</th>
<th>Pst 80% TLC (kPa)</th>
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Mean value
SD
NS
NS
NS
< 0.005
< 0.01
< 0.02
< 0.05

*Pst values at different volumes are given as a percentage of TLC, since their predicted values are most commonly indicated in this way.
†Predicted values according to Yernault et al.8

Discussion

Reduction in expiratory forced flows can be attributed to two main factors, reduction in recoil pressure or an increase in airways resistance. By relating maximal expiratory flow to lung recoil we obtain MFSR curves, as described by Mead et al.8 By means of this curve it is possible to understand the mechanisms of expiratory flow limitation. If the MFSR curve, plotted from these data, were shifted to the right towards the abscissa (see dashed line of figure), it would show that, at any given Pst, MEF would be less than normal. Hence, in this instance, loss of elastic force is not sufficient in itself to explain the low MEF, but increased airways resistance must be responsible. On the other hand, should the decrease of MEF be caused by reduction of elastic recoil, the slope of the MFSR curve would be close to normal or even shifted to the left. It is apparent in this case that although MEF is decreased, it is essentially normal for the available Pst values. In other words, because of the reduction of MEF, the MFSR curve is shorter than normal, but its slope is nearly normal in the absence of bronchial obstruction. Thus, the decreased MEF could be explained by the decreased Pst.

Both loss of recoil and any form of airways obstruction predisposing to airways closure may result in an increase in RV. Under these circumstances RV is determined by the amount of air...
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Figure Maximum flow static recoil (MFSR) curves (---) of 14 subjects showing increased residual volume without bronchial obstruction. Each of the three thick broken lines (-----) represents the mean MFSR curve of five normal subjects, age groups as indicated (derived from Mead et al). The thin dashed line (----) shows the MFSR curve of a patient with severe bronchial obstruction.

trapped in the lungs when the airways are almost all closed. The increase in airways closure at low lung volumes in the elderly is caused mainly by loss of elastic recoil, resulting in airways closure at a higher lung volume than in the young.9

Most of the subjects we studied demonstrated a decrease of MEF 50% and 25% of VC. By means of the MFSR curve it was apparent that this reduction was not the result of increased airways resistance, but rather of loss of elastic recoil. In support of this Pst at 90%, 80%, 70%, and 60% of TLC of the 14 patients showed a significant decrease in comparison to predicted values.

Our results thus demonstrate that in most subjects showing only an increase in residual volume, loss of elastic recoil is the main mechanism. We noticed that while MEF 50% and 25% of VC were significantly less than normal, MEF 75% was not. A possible explanation for this could be that the increase in the cross-sectional diameter of airways at high volumes compensates for the loss of elastic recoil. The effect of these opposing influences is such that MEF at high lung volumes shows little change. At lower volumes, where frictional resistance increases as a function of reduction of the cross-sections of airways, the decrease of MEF, from loss of elastic recoil, becomes more and more evident.

Gelb et al10 studied some patients with a localised lung lesion one week before thoracotomy. Decrease of Pst, a low single breath lung diffusing capacity, and reduced MEF at low lung volumes were found in seven patients. Vital capacity, TLC, FEV1, and Raw, however, were normal or very close to normal. Anatomical studies of lobes or lungs removed within a week of these physiological studies revealed diffuse emphysema in all seven patients.

A displacement to the left of the static pressure-volume curve was observed by Hoepchner et al11 in a group of asymptomatic smokers compared with a control group of non-smokers. On the other hand static compliance was similar in both smokers and non-smokers. This means that measurement of Pst at different lung volumes should be preferred, at least in the diagnosis of incipient emphysema, to the evaluation of static compliance. Of course in the advanced stages of emphysema the pressure-volume curve is displaced to the left, but there is also an
increase in the slope—that is, an increase of static compliance. On the basis of the above data it seems to us that subjects such as ours, with no airways obstruction but with raised RV, probably have emphysema, causing reduction of MEF at medium and low lung volumes as well as the decrease of Pst at different volumes, although static compliance appears to be close to normal.

References