Thoracoabdominal mechanics during tidal breathing in normal subjects and in emphysema and fibrosing alveolitis

NJ BRENNAN, AJR MORRIS, MALCOLM GREEN
From Brompton Hospital, London

ABSTRACT Thoracoabdominal motion and intrathoracic (oesophageal), intra-abdominal (gastric), and transdiaphragmatic pressures were measured during tidal breathing in 20 erect subjects—six normal, seven, with emphysema, and seven with fibrosing alveolitis. In normal subjects all diameters increased synchronously during inspiration and were accompanied by rises in abdominal and transdiaphragmatic pressures. Five patients with fibrosing alveolitis showed a reduction in upper ribcage motion, but normal lower ribcage and abdominal motion; stiff fibrotic lungs may sometimes impair the ability of the diaphragm to expand the upper ribcage and result in proportionally more expansion of the bases. Four emphysematous patients showed increased anteroposterior motion of the ribcage and four showed paradoxical inward motion of the lower ribcage during inspiration. These changes apparently result from increased activity of intercostal and accessory muscles and altered configuration of the diaphragm. The muscles of patients with large, overinflated lungs are at a greater disadvantage than those with small fibroed lungs.

Konno and Mead1 analysed the relative contributions of the ribcage and of the abdomen and diaphragm to ventilation in normal subjects. Several authors have applied similar techniques to patients with severe chronic airways obstruction,2-4 but most groups have studied patients supine and during a variety of contrived respiratory manoeuvres.4-5 Changes in posture from erect to supine affect both static chest wall configuration and its pattern of movement in normal6 and emphysematous subjects.4 We therefore studied normal subjects during tidal breathing in the erect position, since this is the usual posture during daily activities. For comparison we also investigated patients with emphysema (and large lungs) and patients with fibrosing alveolitis (and small lungs). There do not appear to have been any previous studies of patients with fibrotic lung disease.

Methods

The normal subjects were healthy volunteers. Patients in the emphysema group were selected on the basis of clinical history and findings at clinical examination, a compatible chest radiograph, and lung function test values showing hyperinflation with low carbon monoxide transfer coefficient (Kco) (table 1). The patients with fibrosing alveolitis had been diagnosed histologically by lung biopsy. Although the vital capacities of patients with emphysema and fibrosing alveolitis were similar, total lung capacity was nearly twice as great in the emphysema group. Kco was low in both groups. Lung volumes were measured by body plethysmograph and Kco by a helium dilution Resparameter (PK Morgan, Chatham). All patients were ambulatory and they were not in the convalescent phase of an acute infection. Informed consent was obtained.

Table 1 Lung function data for the three groups: mean (±SE) of percentage predicted for each subject

<table>
<thead>
<tr>
<th></th>
<th>Normal (n=6)</th>
<th>Fibrosing alveolitis (n=7)</th>
<th>Emphysema (n=7)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FVC</td>
<td>109±3.6</td>
<td>74±5.2</td>
<td>73±5.9</td>
</tr>
<tr>
<td>FEV1</td>
<td>104±4.8</td>
<td>75±6.8</td>
<td>30±3.4</td>
</tr>
<tr>
<td>TLC</td>
<td>112±6.6</td>
<td>68±4.2</td>
<td>131±6.4</td>
</tr>
<tr>
<td>FRC</td>
<td>118±13.7</td>
<td>66±5.7</td>
<td>165±7.1</td>
</tr>
<tr>
<td>RV</td>
<td>128±13.7</td>
<td>66±5.9</td>
<td>220±14.7</td>
</tr>
<tr>
<td>KCO</td>
<td>102±5.1</td>
<td>61±7.3</td>
<td>43±3.7</td>
</tr>
</tbody>
</table>

FVC—forced vital capacity; FEV1—forced expiratory volume in one second; TLC—total lung capacity; FRC—functional residual capacity; RV—residual volume; KCO—transfer coefficient.

Reprints will not be available.
Thoracoabdominal motion was studied in four diameters with linearised magnetometers attached to the skin with double-surface adhesive tape. Changes in the anteroposterior diameter of the ribcage (R-Ca-p) were measured in the midline 2 cm above the level of the xiphisternal junction, and the a-p diameter of the abdomen (ABDap) in the midline 2 cm above the umbilicus. Changes in ribcage high lateral (R-Chh) and low lateral (R-Chl) diameters were recorded in the mid-axillary line at the level of the xiphisternal junction and the lower costal margin respectively. The R-Chl position was selected because of its proximity to the insertion of the diaphragm. It predominantly spans the abdominal cavity rather than the thorax in normal people. To minimise artefacts due to skin movements, the subjects leaned against a board inclined 17° backwards from vertical, their arms supported away from the chest wall by rests. They were instructed to maintain a constant posture throughout the experiment.

Changes in lung volume were measured at the mouth with an Ohio 840 spirometer. Oesophageal (Poes) and gastric (Pg) pressures were measured with balloons passed into the appropriate positions, standard techniques being used. Both pressures were measured in relation to atmospheric pressure by identical pressure transducers (Mercury M/555). Transdiaphragmatic pressure (Pdi) was derived by electrically subtracting the signal of Poes from Pg and was defined as zero at end-expiratory level during quiet breathing—that is, at functional residual capacity (FRC). Signals from the four magnetometers, volume change, Poes, and Pg were recorded on a magnetic tape multichannel recorder (Racal Store 7) and later displayed against time on a Mingograf 800 recorder or when required on an X-Y recorder. Significance was tested by Student's t test.

Each subject was asked to breathe quietly through the mouthpiece of the spirometer. Four consecutive breaths were analysed after the subjects had become thoroughly accustomed to the apparatus. In each subject means were taken of tidal volume (VT) and diameter change, and the mean of diameter change per litre of inspired volume in each breath was calculated.

Results

Normal subjects

Inspiration from FRC was associated with increase in all chest wall diameters, reaching a maximum at peak tidal inspiration and then returning to the FRC position (fig 1). The relationship between volume and diameter changes was nearly linear and there was little difference between inspiration and expiration (fig 2). During inspiration Poes fell while Pg and Pdi rose; the opposite occurred during expiration (fig 1). Pdi was zero for roughly the second half of expiration.

Fibrosing alveolitis

In patients with fibrosing alveolitis VT did not significantly differ from the values in normal subjects. In three subjects expansion of R-Ca-p and R-Chl was minimal and in two further subjects one of these was substantially reduced. By contrast, R-Chl and ABDap diameters changed normally in all (fig 1). At a given lung volume there were only small differences in chest wall diameters between inspiration and expiration (fig 2). The patterns of changes in Poes, Pr, and Pdi were similar to those in normal subjects (fig 1).

Emphysema

There were appreciable abnormalities of motion in patients with emphysema (fig 3 and table 2). R-Ca-p motion was greater than in normal subjects and much greater than in fibrosing alveolitis. Subjects...
Normal Fibrosing alveolitis

**Fig 2** Change in chest wall diameters against lung volume in two normal subjects (A, B) and two patients with fibrosing alveolitis (C, D). In the fibrosing alveolitis group three patients showed a pattern similar to that of subject C, with reduced R-Cap and R-Ch motion; subject D and the other two subjects showed normal motion. (Abbreviations as in fig 1.)

**Fig 3** Plot of change in chest wall diameter against volume in the seven emphysematous patients to show different patterns. The arrow indicates inspiratory phase. Subject 2 is the same patient as in figure 1. Increased R-Cap motion is seen in subjects 1–3 and paradoxical R-Ch motion in subjects 1–4. Subject 5 shows increased and subject 6 decreased abdominal motion while subject 7 shows relatively normal motions. (Abbreviations as in fig 1.)

**Table 2** Mean values (± SE) for tidal volume (VT) and for increase of chest wall diameters between end-expiration (FRC) and end-inspiration (VT), and chest wall diameters expressed as cm/l of VT for each subject (to allow for differences in VT between subjects)

<table>
<thead>
<tr>
<th></th>
<th>VT (ml)</th>
<th>R-Cap (cm)</th>
<th>R-Ch (cm)</th>
<th>R-Ch-L (cm)</th>
<th>ABD-cap (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>774 ± 223</td>
<td>0.36 ± 0.15</td>
<td>0.30 ± 0.12</td>
<td>0.40 ± 0.20</td>
<td>0.72 ± 0.37</td>
</tr>
<tr>
<td>Fibrosing alveolitis</td>
<td>975 ± 288</td>
<td>0.23 ± 0.13</td>
<td>0.27 ± 0.35</td>
<td>0.66 ± 0.49</td>
<td>0.61 ± 0.56</td>
</tr>
<tr>
<td>Emphysema</td>
<td>1009 ± 290</td>
<td>0.80 ± 0.31**</td>
<td>0.26 ± 0.39</td>
<td>−0.09 ± 0.41*</td>
<td>0.84 ± 0.57</td>
</tr>
</tbody>
</table>

*Abbreviations as in figure 1.

* p < 0.05; **p < 0.01 compared with normal subjects.
(fig 3) showed paradoxical decrease in R-C<sub>1</sub> and in two (subjects 1 and 3) there was also paradoxical change of R-C<sub>b-1</sub> diameter, most noticeable near peak inspiration. Abnormal patterns of ABD<sub>ap</sub> motion were seen in two (subjects 1 and 6), and in three (subjects 2, 3, 5) ABD<sub>ap</sub> diameter was much smaller during expiration than during inspiration at the same lung volume. These abnormalities of chest wall motion did not correlate with lung function values, although the two subjects (1 and 2) with the most evident abnormalities in ribcage motion were the most severely affected by dyspnoea.

During inspiration $P_{oes}$ fell but $P_g$ was variable: it usually fell slightly but in some subjects remained unchanged or rose.

**Discussion**

It has previously been shown that during quiet tidal breathing in erect normal subjects the ribcage and abdominal wall move along their relaxation pressure-volume (p-v) curve. Similarly, the relationship between abdominal pressure and both ribcage and abdominal motion are the same during tidal breathing as during relaxation from total lung capacity. Since the ribcage and abdominal components of the chest wall behave as though relaxed, it has been suggested that the diaphragm is the only important contracting muscle during quiet breathing, and thus minimal energy is needed to expand the chest wall. Our results in normal subjects are compatible with this analysis. In fibrosing alveolitis, however, we found that movement of the upper ribcage was reduced in five out of seven patients. In emphysema the patterns were varied and disorganised.

In fibrosing alveolitis the reduced lung volume tends to cause the diaphragm to be lengthened and highly curved. This would place the muscle at a mechanical advantage. Recent work has shown that diaphragmatic activity produces greater pleural pressure swings near the bases than at the lung apices in dogs, and there is evidence of similar pressure gradients in normal man, which may affect ventilation distribution. The decreased pulmonary compliance in fibrosing alveolitis may exacerbate these pressure gradients down the lungs and accentuate the normal top-to-bottom ventilation gradient. Preferential ventilation of the lung bases could explain the clinical observation that crepitations are maximal in this zone.

It is well recognised that in emphysema there is increased activity of the intercostals and accessory muscles of respiration, and consequently these patients show a relatively larger contribution of ribcage displacement to ventilation than normal subjects. Such activity presumably distorts the chest wall from its relaxation p-v curve, thus increasing the work of breathing. Furthermore, abnormal patterns of abdominal wall motion have been described in patients with chronic airways obstruction, particularly when supine. Our emphysematous patients did show features compatible with excessive activity of intercostal and accessory muscles during inspiration: increased amplitude of R-C<sub>ap</sub> excursions and little change or a fall in $P_g$. The latter is presumably due to ribcage elevation by muscles other than the diaphragm. In addition, in some individuals ABD<sub>ap</sub> diameter during expiration was smaller than during inspiration at the same lung volume. This implies activity of abdominal muscles during expiration. Conceivably such activity may aid diaphragmatic function by exerting an upward force to increase diaphragm length and curvature before the next inspiration.

Some of our emphysematous patients had paradoxical inward motion of the lower ribcage during inspiration. Clinically this has been called Hoover's sign and is thought to result from inward tension exerted by the flattened diaphragm. In addition, the flat diaphragm exposes the lower ribcage to negative intrapleural pressure generated by both diaphragm and the overactive intercostal and accessory muscles, whereas in normal subjects the presence of an area of apposition between diaphragm and ribcage means that the lower ribcage is effectively exposed to a positive outward, intra-abdominal pressure during inspiration. The mobility of these lowermost "floating" ribs may cause them to deform easily in response to such changes in the forces acting on them.

We conclude that emphysema leads to disorganised and apparently inefficient movement, even during tidal breathing. Whether training such patients to breathe with a pattern more similar to that of normal people is practicable or useful remains to be established, perhaps by studies such as these before and after training. In fibrosing alveolitis expansion of the upper part of the lungs appears to be impaired despite an increase in the mechanical advantage of the diaphragm. This may be due to the stiffness of the lungs and might explain basal crepitations and, in part, the reduction of gas transfer. Conceivably muscle training in these subjects should be directed, if at all, to their upper ribcages.

**References**


