Rib cage mechanics after median sternotomy

T J Locke, T L Griffiths, H Mould, G J Gibson

Abstract
A substantial reduction in lung volumes occurs after sternotomy, but the mechanism or mechanisms are unclear. Measurements were made of lung volumes and of chest wall motion with four pairs of magnetometers (two pairs for anteroposterior rib cage, one for lateral rib cage, and one for anteroposterior abdominal dimensions) in 16 men before and one week and three months after coronary artery grafting. Reductions in all lung volumes occurred after sternotomy and were greater in the supine than in the sitting position. Supine vital capacity was reduced one week after surgery, with almost complete recovery at three months. One week after sternotomy there was a significant reduction in tidal volume from a mean (95% confidence limits) value of 0.88 (0.76–1.00) litre to 0.61 (0.32–0.70) l, and in supine rib cage displacement from 3.87 (1.96–5.78) mm to 0.44 (–0.61 – 1.49) mm in the lateral plane. Respiratory frequency increased from 16 (13–19) to 21 (19–24)/min. Coordination of the rib cage was assessed by measuring the difference in timing of onset of chest wall motion and airflow in four planes. At one week nine of 14 patients showed uncoordination between airflow and rib cage motion in one or more dimensions, and this was still present in three patients at three months. No loss of the temporal relation between airflow and abdominal wall motion was detected. The results suggest that reduced and uncoordinated rib cage expansion contributes to the restrictive ventilatory defect that follows median sternotomy.

MEASUREMENTS
FEV1 and slow expired vital capacity (VC) were measured in both the upright and supine positions and static lung volumes (total lung capacity (TLC), functional residual capacity (FRC), and residual volume (RV)) were measured with the subject seated in a whole body plethysmograph. In addition, displacement of the chest wall was measured during resting tidal breathing with four pairs of linearised magnetometers, attached to the skin by double sided tape in the positions shown in figure 1. Signals were recorded on a multichannel pen recorder (Lectromed M19). The linear displacement of each pair of magnetometers was calibrated by a standard electrical signal. The patient breathed with a mouthpiece and nose clip through a Fleisch...
pneumotachograph to allow accurate identification of the phases of respiration, and the output from the pneumotachograph was integrated electrically to give tidal volume.

In normal subjects during tidal breathing the chest wall moves symmetrically with linear and volume expansion of its rib cage and abdominal compartments closely related in time. When chest wall expansion is distorted differences in phase arise between motion in different dimensions. The distortions are conveniently quantified by assuming that volume displacement and linear motion approximate to a sine wave and by measuring the "phase angle" between the two signals (fig 2). Although chest wall expansion does not strictly follow a sine wave pattern, a similar approach has been applied successfully in quantifying rib cage distortion in severe asthma. In the present study phase angles between linear and volume displacement were measured for each dimension in each of a sequence of five breaths on each occasion the patient was studied. In addition, the amplitudes of volume and linear displacements from end expiration to end inspiration were calculated during a further 60 breaths. The net linear displacement defined in this way is inevitably affected by the phase angle; in the example shown in figure 2 delay in linear motion (positive phase angle) implies reduced expansion in this dimension between end expiration and end inspiration. In the most extreme case dimensional change would be of opposite sign to volume change throughout the cycle and the corresponding "phase angle" would be 180°.

Baseline ranges for phase angles for each dimension were established from the preoperative study and postoperative values outside this range of mean and two standard deviations were regarded as abnormal.

In eight patients respiratory muscle function was assessed by recording maximum static inspiratory (Pimax) and expiratory (Pemax) pressures at the mouth. Expiratory efforts were initiated at TLC and inspiratory efforts at FRC; the values recorded were sustained for at least one second and represented the best of three efforts.

Patients were studied before and one week and three months after operation. Two patients declined magnetometer measurements at one week. In all cases a chest radiograph was available within two days of the first postoperative study day; radiographs were not taken at the time of the three month study.

STATISTICS
In the text and figures data are presented as means with 95% confidence limits in parentheses. Results before and after surgery were compared by means of the paired t test or the Spearman correlation coefficient (r) as appropriate. The number of patients with abnormal phase angles after operation was compared with the number before operation with Fisher's exact test.

Results
Postoperative respiratory complications evident one week after sternotomy were minor; plate atelectasis was visible radiographically in four patients, and small pleural effusions were present in six; three patients were being treated for acute bronchitis. No patient had radiographic features suggesting phrenic nerve injury. One patient had a left lower brachial plexus injury. There were no instances of sternal dehiscence.

The changes in lung volumes one week and three months after sternotomy are shown in table 1. As expected, a restrictive ventilatory defect occurred and this had largely recovered at three months. The fall in FEV1 in six patients who had internal mammary grafts (from 3.13 (95% CL 2.69–3.57) to 1.60 (1.24–2.06) l) was similar to that in patients who had only saphenous vein grafts (3.09 (2.55–3.63) to 1.65 (1.26–2.04) l). Reductions in lung volumes one week...
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<table>
<thead>
<tr>
<th>Table 1</th>
<th>Lung volumes in the erect posture before and after sternotomy (mean (95% confidence limits))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
</tr>
<tr>
<td>FEV (l min⁻¹)</td>
<td>3.40 (3.08-3.72)</td>
</tr>
<tr>
<td>VC (l)</td>
<td>4.38 (4.03-4.73)</td>
</tr>
<tr>
<td>TLC (l)</td>
<td>7.56 (6.96-8.16)</td>
</tr>
<tr>
<td>FRC (l)</td>
<td>4.72 (4.16-5.28)</td>
</tr>
<tr>
<td>RV (l)</td>
<td>3.29 (2.95-3.65)</td>
</tr>
</tbody>
</table>

Values of p (*p < 0.01, tp < 0.02) were derived from paired t tests of control and postoperative data.

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Rib cage displacements during tidal breathing in the supine posture (mean (95% confidence limits))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
</tr>
<tr>
<td>Manubrium (mm)</td>
<td>1.74 (0.91-2.57)</td>
</tr>
<tr>
<td>Xiphoid (mm)</td>
<td>2.70 (1.57-3.84)</td>
</tr>
<tr>
<td>Abdomen (mm)</td>
<td>13.3 (10.9-15.8)</td>
</tr>
<tr>
<td>Lateral (mm)</td>
<td>3.87 (1.96-5.78)</td>
</tr>
<tr>
<td>Displacement rate</td>
<td>16 (13-19)</td>
</tr>
<tr>
<td>VT (l)</td>
<td>0.88 (0.76-1.00)</td>
</tr>
</tbody>
</table>

Values of p (*p < 0.05, tp < 0.01, **p < 0.001) were derived from paired t tests of control and postoperative data.

Maximal expiratory pressure fell significantly; there was a slight reduction in maximum inspiratory pressure, but this was not significant (table 4).

Discussion

Restrictive ventilatory defects after sternotomy were recognised in 1962, but the mechanisms have not been completely elucidated. The changes are most severe in the first 48 hours after operation. By one week after surgery some recovery has occurred and by three months recovery is almost complete. Our values lie within the range of previous results, though in our relatively small group of patients we did not find the more substantial reduction in lung volumes associated with the use of the internal mammary artery reported by others.

In the early stages after coronary artery bypass grafting pain atelectasis and the transient increase in lung water that follows cardiopulmonary bypass would account for a major portion of the fall in VC. By one week, however, all the patients in the present study were ambulant and no longer required opiate analgesics; no patient complained of pain during the forced manoeuvres. Postoperative chest radiographs excluded obvious causes of major volume loss, such as lobar collapse, consolidation, or large pleural effusion. Occult pulmonary oedema cannot be excluded completely but is unlikely to contribute to an important extent to the reduction in lung volume seen one week after surgery; no patient had dyspnoea or radiographic evidence of pulmonary oedema. Fluid that accumulates in the lungs after car-

<table>
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<tr>
<th>Table 3</th>
<th>Number of patients with abnormal phase angles</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Control</td>
</tr>
<tr>
<td>Manubrium</td>
<td>0</td>
</tr>
<tr>
<td>Xiphoid</td>
<td>0</td>
</tr>
<tr>
<td>Abdomen</td>
<td>0</td>
</tr>
<tr>
<td>Lateral</td>
<td>0</td>
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</tbody>
</table>

*p = 0.04, **p = 0.0008 (Fisher's exact test).
Table 4  Maximum respiratory pressures (mean (95% confidence limits))

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>One week</th>
<th>Three months</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pmax (cmH₂O)</td>
<td>150 (109-191)</td>
<td>80 (50-109)</td>
<td>113 (70-156)</td>
</tr>
<tr>
<td>Pimax (cmH₂O)</td>
<td>100 (49-150)</td>
<td>83 (44-123)</td>
<td>80 (38-122)</td>
</tr>
</tbody>
</table>

The value of p (*p < 0.01) was derived from a paired t test of control and postoperative data.

Pmax—maximum expiratory pressure; Pimax—maximum inspiratory pressure.

diopulmonary bypass has usually cleared by 48 hours after operation,13,14 and there is little difference in lung compliance before and 8–13 days after coronary artery grafting.15

Estenne et al4 investigated diaphragmatic function postoperatively and showed normal phrenic nerve conduction times in 11 of 12 patients after saphenous vein grafting, suggesting that phrenic injury is unlikely to contribute to volume reduction in most patients. In the present study indirect indices of diaphragmatic function (Pmax, abdominal movement phase angle, and chest radiograph) support the conclusion that weakness of the diaphragm was not a factor in the postoperative change in pulmonary function. The small but significant decrease in supine abdominal wall displacement during tidal breathing at one week might be explained by the extension of the sternotomy into the upper abdomen, producing reflex inhibition of motion.16 Similar factors may account for the impressive reduction in Pmax, which presumably reflects impaired function of the abdominal wall muscles.

Pleurotomy seems unlikely to be relevant to the abnormalities found: mobilisation of the left internal mammary artery was performed without pleurotomy if possible, and the pleural space was entered in only two patients; furthermore, the patients who had only saphenous vein grafting showed changes similar to those of the ones who also had internal mammary artery grafting. No patient had an intercostal drain, which may increase respiratory abnormalities.4,5

The possibility that alterations of rib cage mechanics14 play a part in the reduction of lung volumes is suggested by the generalised decrease in rib cage motion and the relation between the decrease in vital capacity and the reduction of lateral rib cage motion. Two patterns of abnormality of chest wall movement were identified. Five patients had a simple reduction of displacement in one or more planes without any significant changes in the timing of the onset of chest wall motion; the remainder had abnormalities of timing of chest wall motion and consequently reduced displacements. The presence of frank paradox of the lateral rib cage dimension in four patients represents an extreme form of this type of discoordination. A possible mechanism may be trauma to the costovertebral joints, producing reflex inhibition of intercostal muscle contraction such that the rib cage is unable to resist the upward force of diaphragmatic contraction.17 There may also be a change in the bony configuration of the chest wall after sternotomy with a increase in thoracic spinal curvature and lowering of the ribs.18

We conclude that the restrictive ventilatory defect that follows median sternotomy is likely to be due in part to alterations in rib cage mechanics.

We thank Mr AH Brown and Mr CGA McGregor for permission to study their patients.

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