Respiratory mechanics in patients with tense cirrhotic ascites


Eur Respir J 1997; 10: 1622–1630

ABSTRACT: Lung volumes are decreased by tense ascites and increase after large volume paracentesis (LVP). The overall effect of ascites and LVP on the respiratory function is poorly understood.

We studied eight cirrhotic patients with tense ascites before and after LVP. Inspiratory muscle force (maximal transdiaphragmatic pressure (Pdi,max)), and the lowest pleural pressure (Ppl,min)) was assessed while the patients were seated. Rib cage and abdominal volume displacements, as well as pleural and gastric pressures were measured during quiet breathing while the patients were supine.

Pdi,max and Ppl,min were normal and did not change after LVP (from 84.2±19.7 to 85.2±17.0 cmH2O and from 68.3±19.7 to 74±15.9 cmH2O, respectively). The abdominal contribution to the generation of tidal volume was greater than that of the rib cage (79 vs 21%), a pattern which did not change after LVP (73 and 27%).

Before LVP, tidal swings both of pleural pressure (Ppl,sw) and transdiaphragmatic pressure (Pdi,sw) were large (15.3±4.3 and 18.5±3.9 cmH2O, respectively) and the load on inspiratory muscles was increased as a consequence of elevated dynamic elastance of the lung (Edyn) (11.4±2.6 cmH2O·L-1) and ("atmospheric") positive end-expiratory pressure (PEEPi) (4.3±3.5 cmH2O). LVP reduced the load on the inspiratory muscles, as shown by the significant decrease in Ppl,sw (10.6±2.0 cmH2O), Pdi,sw (12.8±3.0 cmH2O), Edyn (10.0±2.0 cmH2O·L-1) and PEEPi (1±1.3 cmH2O).

The amount of fluid removed was closely related to changes in Ppl,sw and PEEPi.

We conclude that the strength of the inspiratory muscles is normal or reduced in seated cirrhotic patients. In the supine position, tense ascites results in an increase in lung elastic load and development of positive end-expiratory pressure, with a consequent overload and increased activation of inspiratory muscles. Large volume paracentesis decreases overloading and activation, but does not change the strength of the inspiratory muscles.


Although tense ascites has been reported to decrease lung volumes, and large volume paracentesis (LVP) to increase them [1–3], the overall effect of ascites and LVP on the respiratory function is poorly understood. In patients with ascites, ABELMANN et al. [1] observed decreased lung volumes, rapid shallow breathing and increased resting oxygen consumption; paracentesis caused an increase in lung volumes, and a reduction both of respiratory frequency and resting oxygen consumption. According to ABELMANN et al. [1], these alterations were due to the increased intra-abdominal pressure that was transmitted to the chest, causing increase in pleural pressure, elevation and relative fixation of the diaphragm, and increased stiffness of the chest wall: the movements of a more rigid thoracic cage required an increased work of breathing. If this hypothesis is correct, overloading and, thereby, increased activation of inspiratory muscles should be observed in ascitic patients. These aspects of the respiratory function have not been defined [2–4]. To obtain further insight into this issue, we carried out the present study, in which the mechanical characteristics of the respiratory system were investigated in patients with tense ascites before and after LVP.

Methods

Subjects

Eight patients (7 males and 1 female; mean age 57±8 yrs, range 43–71 yrs) with tense cirrhotic ascites, consecutively admitted to the Istituto di Medicina Interna of the University of Florence between October 1993 and March 1995, were studied. Tense ascites was defined as "a tight distension of abdominal wall consequent to the presence of a high intra-abdominal pressure in cirrhotic patients with large ascitic fluid accumulation evoking a strong sense of abdominal painful tension". Criteria for inclusion in the study were: 1) presence of tense ascites; 2) no clinical or laboratory evidence of
other systemic diseases, infection or fever, recent gastrointestinal bleeding, grade III or IV encephalopathy [5], hepatocellular carcinoma; 3) no history of alcoholism; 4) no parenchymal lung involvement or pleural effusion on routine chest radiographic image; and 5) absence of diagnostic criteria for asthma, chronic bronchitis or emphysema according to the American Thoracic Society [6]. The anthropometric and clinical data of the patients are presented in table 1. Two patients were non-smokers, and the remaining six former smokers. Cirrhosis had been diagnosed by history, clinical examination, laboratory findings and, when not contra-indicated, liver biopsy. Viral hepatitis was the cause of cirrhosis in seven patients, and cryptogenic cirrhosis was diagnosed in one patient. From a clinical point of view, the patients' conditions varied from moderate to severe according to the Child-Pugh classification [7]. None of the patients complained of dyspnoea.

After admission to hospital, patients were given a diet containing 40 mEq·day⁻¹ of sodium, with a controlled fluid intake (≤1 L·day⁻¹). In patients with hyponatraemia (serum sodium <130 mEq·L⁻¹) water ingestion was restricted to 500 mL·day⁻¹.

The protocol of the study was approved by the University's Ethics Committee and written informed consent was obtained from each subject.

**Protocol**

The study was conducted on three consecutive days. On the first day, blood gas values, lung volumes, and inspiratory muscle strength were measured, while the patients were seated in a comfortable high-backed armchair; they were then placed in a comfortable supine position and breathed quietly through the pneumotachograph. After a 10 min period of acclimatization, breathing pattern variables, rib cage and abdominal volume displacements, and pleural and gastric pressure swings were recorded during tidal breathing. On the second day, large volume paracentesis (LVP) was performed. LVP was defined as removal of >2,000 mL of ascitic fluid at a single tap. The amount of fluid removed with paracentesis was variable, but always >3.5 L. Paracentesis was performed via gravity with a No. 16 gauge catheter, under local anaesthesia, in the left lower abdominal quadrant over a 5 h period, using standard aseptic technique. Patients were initially in the supine position; they were turned into a lateral position if necessary. Ultrasound was performed to define the best position for the patient, in order to remove the largest quantity of ascites.

After tap, patients reclined for 2 h on the opposite side to prevent leakage of ascitic fluid. Samples of ascitic fluid were taken during each paracentesis for total and differential white blood cell counts, cytology, biochemical examination and cultures. During each paracentesis, human albumin (human albumin 20%; Immuno AG, Austria) was administered intravenously at a dosage of 10 g per litre of ascitic fluid removed [8]. On the third day, pulmonary function testing was repeated, using the same methods as employed on the first day. On the second and third day, blood electrolytes and body weight were recorded.

**Measurements**

Routine spirometry was performed using a water-sealed spirometer (Pulmonary Godart; Sensormedics Corp., Yorba Linda, CA, USA). Functional residual capacity (FRC) was measured by the helium dilution technique. The normal values for lung volumes were those of the European Coal and Steel Community [9]. Arterial blood samples were taken while the subjects were breathing room air, and blood gas values were analysed (ABL-3 analyzer; Radiometer, Copenhagen, Denmark).

For ventilation measurements, patients breathed through a Fleisch No. 3 pneumotachograph connected to a differential pressure transducer. Volume was obtained by electrical integration of the flow signal. From the spirogram the following parameters were derived: inspiratory time (ti), expiratory time (te), total time of the respiratory cycle (tot) and tidal volume (VT). Respiratory frequency (r = 1/tot × 60) and minute ventilation (VE = VT × r) were also calculated. End-tidal carbon dioxide tension (PETCO₂) and arterial oxygen saturation (Sao₂) were monitored continuously by an infra-red CO₂ meter (Datex Normocap, Helsinki, Finland) and an ear oximeter (Radiometer, Copenhagen, Denmark), respectively.

Mouth pressure during tidal breathing was measured using a pressure transducer (Statham P23ID; Statham Lab. Inc., Hato Rey, Puerto Rico).

Changes in thoracoabdominal dimensions were determined in 6 of the 8 patients by linearized magnetometers. Pairs of magnetometer coils were attached on the midline to measure the anteroposterior diameters of the lower rib cage at the level of the nipples (fifth costal cartilage) and of the abdomen approximately 2 cm above the umbilicus. The magnetometer output voltages were displayed on an x-y Tektronics 5115 storage oscilloscope (Tektronics Corp., Beaverton, OR, USA), and photographed using a Polaroid camera (Polaroid Corp.,

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**Table 1. – Anthropometric and clinical data of patients with cirrhotic ascites**

<table>
<thead>
<tr>
<th>No.</th>
<th>Sex</th>
<th>Age yrs</th>
<th>Weight kg</th>
<th>Height cm</th>
<th>Smoking pack-yrs</th>
<th>Diagnosis</th>
<th>Child-Pugh class</th>
<th>Amount of fluid removed L</th>
<th>Weight decrease kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>M</td>
<td>71</td>
<td>76</td>
<td>168</td>
<td>40</td>
<td>HBV</td>
<td>B</td>
<td>11.2</td>
<td>11.0</td>
</tr>
<tr>
<td>2</td>
<td>M</td>
<td>43</td>
<td>87</td>
<td>174</td>
<td>17</td>
<td>HBV, HCV</td>
<td>C</td>
<td>5.1</td>
<td>5.5</td>
</tr>
<tr>
<td>3</td>
<td>M</td>
<td>59</td>
<td>92</td>
<td>166</td>
<td>10</td>
<td>CRYPT</td>
<td>C</td>
<td>13.0</td>
<td>15.1</td>
</tr>
<tr>
<td>4</td>
<td>M</td>
<td>66</td>
<td>74</td>
<td>164</td>
<td>24</td>
<td>HCV</td>
<td>C</td>
<td>6.8</td>
<td>6.7</td>
</tr>
<tr>
<td>5</td>
<td>M</td>
<td>55</td>
<td>85</td>
<td>170</td>
<td>0</td>
<td>HBV</td>
<td>C</td>
<td>6.5</td>
<td>7.0</td>
</tr>
<tr>
<td>6</td>
<td>F</td>
<td>56</td>
<td>62</td>
<td>150</td>
<td>0</td>
<td>HCV</td>
<td>C</td>
<td>4.1</td>
<td>6.0</td>
</tr>
<tr>
<td>7</td>
<td>M</td>
<td>53</td>
<td>106</td>
<td>183</td>
<td>36</td>
<td>HCV</td>
<td>C</td>
<td>3.5</td>
<td>5.5</td>
</tr>
<tr>
<td>8</td>
<td>M</td>
<td>55</td>
<td>72</td>
<td>160</td>
<td>45</td>
<td>HCV</td>
<td>C</td>
<td>7.2</td>
<td>9.0</td>
</tr>
</tbody>
</table>

Pts: patients; M: male; F: female; HBV: hepatitis B virus; HCV: hepatitis C virus; CRYPT: cryptogenic cirrhosis.
Cambridge, MA, USA). Signal gains were adjusted so that isovolume manoeuvres ("belly-in") produced a slope of approximately -1 when the anteroposterior diameter of the abdomen was plotted against the anteroposterior diameter of the lower rib cage [10].

Oesophageal pressure (Poes) was measured with a conventional balloon-tipped catheter system [11] connected to a differential pressure transducer (Validyne, Northridge, CA, USA), as described previously [12]. The balloon was positioned in the mid-oesophagus and contained 0.4 mL of air. Oesophageal pressure was used as an index of pleural pressure (Ppl). Gastric pressure (Pga) was simultaneously measured with a similar balloon-tipped catheter system connected to a second differential pressure transducer. This balloon was positioned in the stomach with the tip 65–70 cm from the nares, and contained 2 mL of air. Transdiaphragmatic pressure (Pdi) was obtained by subtracting Ppl from Pga.

Inspiratory muscle strength was assessed by measuring minimal (i.e. the greatest negative) inspiratory pleural pressure (Ppl,min) and maximal transdiaphragmatic pressure (Pdi,max) at FRC during sniff manoeuvres [13]. The patients were repeatedly encouraged to try as hard as possible, and they had a visual feedback of the pressure generated. The manoeuvres were repeated until three measurements with less than 5% variability were recorded. The lowest Ppl,min and the highest Pdi,max values obtained were used for analysis and compared with those of age-matched normal subjects: the upper limit of normality was calculated as the mean value +SD×1.65.

Ppl and Pga were also recorded during tidal breathing: Ppl (Ppl,sw) and Pga (Pga,sw) swings were calculated as the difference between the pressure measured at end-expiration and the peak value measured during inspiration. These values were compared with those of age-matched normal subjects: the upper limit of normality was calculated as the mean value +SD×1.65. Total lung resistance (RL) and lung dynamic elastance (Eldyn) were measured during breathing at rest. Total lung resistance was obtained using the isovolume method of Frank et al. [14]. Dynamic lung elastance was determined by dividing the difference in Ppl between points of zero flow by VT. To evaluate end-expiratory alveolar pressure, we used the indirect method recently described by Haluzska et al. [15] and Dal Vecchio et al. [16], rather than the direct method of airway occlusion. This was because awake subjects react to airway occlusion in an unpredictable fashion, so that no reliable measurements of alveolar pressure can be obtained. We thus looked for the presence of a time lag between the fall in Ppl at the onset of the inspiratory effort and the onset of inspiratory airflow, and measured the negative deflection in Ppl that preceded the start of inspiratory flow (fig. 1). This negative deflection in Ppl will be referred to as intrinsic positive end-expiratory pressure (PEEPi) for consistency with previous investigations [15, 16].

The change in Pga resulting from the contraction of the abdominal muscles during expiration was also assessed. In agreement with Ninane et al. [17], the increase in Pga during the expiratory phase of the breathing cycle was taken as a reflection of the mechanical effect of abdominal muscle contraction. The ratio of pleural pressure swing to tidal volume (Ppl,sw/VT) was also calculated, in order to assess the pressure necessary to produce tidal volume.

All signals were recorded continuously on a multichannel chart recorder (TA4000; Gould, Valley View, OH, USA).

**Statistical analysis**

Mean values and standard deviations of the mean were calculated for all variables. Data obtained under control conditions and after LVP were compared by Wilcoxon test for paired samples. Single and stepwise multiple regression analyses were performed to assess the relationships between variables. The proportion of total variance of the dependent variable accounted for by the predictor variable(s) is reported as the square of correlation coefficient (r²), expressed as a percentage. Single regression analysis was carried out by least square method. All statistical analyses were carried out using the SPSS for Windows 6.0 package (SPSS Inc., Chicago, IL, USA).

**Results**

In all patients, the decrease in body weight from the second to the third day was equal to or greater than the weight of fluid removed; moreover, there was a close relationship between the amount of fluid removed and the decrease in body weight (r=0.96; p<0.0002). Blood electrolytes did not significantly change after LVP.
Lung volumes

Pulmonary function data before and after LVP are presented in table 2. Before LVP, vital capacity (VC), FRC, and forced expiratory volume in one second (FEV₁) varied from normal to moderately reduced; total lung capacity (TLC) and residual volume (RV) were normal in all but one subject, while FEV₁/VC was slightly reduced in three subjects. Expiratory reserve volume (ERV) was markedly reduced in four subjects and slightly reduced in three. After LVP, the FRC, FEV₁ and ERV increased significantly, while inspiratory capacity (IC) decreased significantly; VC also increased but the change did not reach the level of significance, while TLC, RV and FEV₁/VC remained largely unchanged.

Breathing pattern and blood gas values

Before LVP, V'E was elevated (14.4±2.4 L·min⁻¹), due to large tidal volume (0.82±0.23 L) and high respiratory frequency (18.5±4.3 breaths·min⁻¹). No changes were observed after LVP (V'E = 13.7±1.8; V'T = 0.81±0.25 L; f'R = 17.8±3.8 breaths·min⁻¹). Mean pH was within normal limits, arterial oxygen tension (Pₐ,O₂) was only slightly reduced and arterial carbon dioxide tension (Pₐ,CO₂) was low; LVP did not cause significant changes (table 2).

Table 2. – Pulmonary function data, arterial blood gas values and mechanical lung characteristics in the patients with cirrhotic ascites

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Before LVP</th>
<th>After LVP</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>VC % pred</td>
<td>90±10</td>
<td>98±17</td>
<td>NS</td>
</tr>
<tr>
<td>FRC % pred</td>
<td>86±19</td>
<td>98±14</td>
<td>&lt;0.014</td>
</tr>
<tr>
<td>TLC % pred</td>
<td>94±10</td>
<td>97±11</td>
<td>NS</td>
</tr>
<tr>
<td>RV % pred</td>
<td>107±21</td>
<td>103±24</td>
<td>NS</td>
</tr>
<tr>
<td>FEV₁ % pred</td>
<td>87±12</td>
<td>100±17</td>
<td>&lt;0.009</td>
</tr>
<tr>
<td>FEV₁/VC %</td>
<td>74±7</td>
<td>79±5</td>
<td>NS</td>
</tr>
<tr>
<td>IC L</td>
<td>2.9±0.63</td>
<td>2.7±0.53</td>
<td>&lt;0.029</td>
</tr>
<tr>
<td>ERV L</td>
<td>0.58±0.30</td>
<td>0.98±0.45</td>
<td>&lt;0.014</td>
</tr>
<tr>
<td>Pₐ,O₂ mmHg</td>
<td>11.96±1.03</td>
<td>12.12±0.64</td>
<td>NS</td>
</tr>
<tr>
<td>Pₐ,CO₂ mmHg</td>
<td>4.37±0.43</td>
<td>4.52±0.36</td>
<td>NS</td>
</tr>
<tr>
<td>pH</td>
<td>7.44±0.02</td>
<td>7.44±0.02</td>
<td>NS</td>
</tr>
<tr>
<td>Rₜ cmH₂O·L⁻¹·s⁻¹</td>
<td>5.9±1.6</td>
<td>5.0±1.5</td>
<td>NS</td>
</tr>
<tr>
<td>Eᵋ dyn cmH₂O·L⁻¹·s⁻¹⁻¹</td>
<td>11.4±2.6</td>
<td>10.0±2.0</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>PEEP cmH₂O</td>
<td>4.3±3.5</td>
<td>1.1±1.3</td>
<td>&lt;0.02</td>
</tr>
</tbody>
</table>

Values are presented as mean± SD. VC: vital capacity; FRC: functional residual capacity; TLC: total lung capacity; RV: residual volume; FEV₁: forced expiratory volume in one second; IC: inspiratory capacity; ERV: expiratory reserve volume; Pₐ,O₂: arterial partial tension of oxygen; Pₐ,CO₂: arterial partial tension of carbon dioxide; Rₜ: total lung resistance; Eᵋ dyn: dynamic elastance of the lung; PEEP: positive end-expiratory alveolar pressure; % pred: percentage of predicted value; LVP: large volume paracentesis; NS: nonsignificant. (1 mmHg= 0.133 kPa).

Fig. 2. – a) Minimal pleural pressure (Pₚₗ₉ₘᵋₗ₉ₜₗ₉ₑ₉) in cmH₂O; b) maximal transdiaphragmatic pressure (Pₚₛₐₘ₉₉ₚₑ₉ₔ) in cmH₂O; c) pleural pressure swings (Pₚₙₙ₉ₘ₉ₛ₉₉ₚₑ₉ₔ) in cmH₂O; and d) transdiaphragmatic pressure swings (Pₚₚₚₚ₉ₐ₉ₚₑ₉ₔ₉ₙ₉ₚₑ₉ₔ,₉ₐ₉ₚₑ₉ₔ) in cmH₂O before and after large volume paracentesis (LVP). Dashed lines represent the lower limits of the normal range (mean -1.65 SD) for males (M) and females (F), respectively. ●: male; ○: female; NS: nonsignificant.
**Inspiratory muscle strength**

The mean inspiratory muscle strength in the sitting position was not significantly different from that of normal subjects. However, looking at individual values, before LVP, \(P_{pl, min}\) was normal in four patients, just below the lower normal limit in two, and low in the remaining two (fig. 2a), whereas \(P_{di, max}\) was low in six patients and normal in two (fig. 2b). After LVP, neither \(P_{pl, min}\) nor \(P_{di, max}\) changed consistently (fig. 2a and b).

**Chest wall mechanics**

Compared to the normal range of our laboratory, swings of pleural pressure (\(P_{pl, sw}\)) (figs. 2c and 3) and transdiaphragmatic pressure (\(P_{di, sw}\)) (figs. 2d and 3) during tidal breathing in the supine position were large before LVP and fell significantly thereafter. The gastric pressure swing (\(P_{ga, sw}\)) also decreased significantly with LVP (from 3.2 to 2.1 cmH\(_2\)O; \(p<0.01\)). Plots of \(P_{pl}\) versus \(P_{ga}\) before and after LVP are illustrated in figure 3. It appears that with LVP: 1) both end-expiratory pleural pressure (\(P_{pl, EE}\)) and end-expiratory gastric pressure (\(P_{ga, EE}\)) decreased significantly (\(p<0.02\) for both); and 2) end-inspiratory pleural pressure (\(P_{pl, EI}\)) did not change, whilst end-inspiratory gastric pressure (\(P_{ga, EI}\)) decreased markedly (\(p<0.01\)).

The magnetometry data show that during quiet breathing in supine position the rib cage contributed 0.18 L and abdomen 0.72 L (21 and 79%, respectively) to tidal volume and these contributions did not change significantly after LVP (rib cage 0.21 L and abdomen 0.64 L (27 and 73%, respectively)) (fig. 4a). The ratio of abdominal displacement (\(V_{ab}\)) to \(P_{ga, sw}\) (\(V_{ab}/P_{ga, sw}\)) during quiet breathing was not significantly modified by LVP (0.238 and 0.295 L·cmH\(_2\)O\(^{-1}\) before and after LVP, respectively). In four patients, the changes \(V_{ab}/P_{ga, sw}\) were minor, whilst in the other two patients \(V_{ab}/P_{ga, sw}\) increased markedly. Plots of \(V_{ab}/P_{ga, sw}\) in a representative subject before and after LVP are presented in figure 4b.

**Lung mechanics**

In the supine position, \(R_t\) was slightly increased [9]. \(E_l,\text{dyn}\) was high in all subjects and PEEPi was present in 7 out of 8 patients. After LVP, \(R_t\) did not change significantly, whereas both \(E_l,\text{dyn}\) and PEEPi decreased significantly (table 2 and fig. 1).

**Interrelated measurements**

Changes in \(P_{pl, EE}\), \(P_{pl, sw}\) and PEEPi were closely related to the amount of fluid removed (fig. 5a–c). Moreover, changes in PEEPi with LVP were significantly related to changes in \(P_{pl, sw}/VT\) induced by LVP were closely related to changes in \(E_l,\text{dyn}\) and PEEPi (\(r=0.83;\) \(p<0.02\); and \(r=0.91;\) \(p<0.002\), respectively). Stepwise multiple regression analysis (table 3) showed that changes in \(E_l,\text{dyn}\) (\(F=7.32;\) \(p=0.042\)) and PEEPi (\(F=18. 14;\) \(p=0.008\)) induced by LVP explained 90% of the variability of changes in \(P_{pl, sw}/VT\). There were no significant relationships between PEEPi and FEV\(_1\), FEV\(_1\)/FVC, or breathing pattern.
The main findings of the present study can be summarized as follows. Firstly, inspiratory muscle strength, measured in a sitting position, was normal or reduced and was not modified by LVP. Secondly, in the supine position, pleural and transdiaphragmatic pressure swings during tidal breathing were large and decreased with LVP; the lung elastic load was increased and a threshold load was present in most patients; both elastic and threshold load decreased with LVP.

A limitation of this study is represented by the fact that no ultrasound examination was performed on the third day in order to exclude the possibility that ascites developed after LVP. However, the following points should be considered: 1) the decrease in body weight was equal to or greater than the weight of fluid removed in all patients, and there was a close relationship between the two; and 2) during paracentesis, patients were given large amounts of human albumin (see Methods), which has been shown [18] to prevent the occurrence of hypovolaemia and, therefore, rapid reaccumulation of ascites. We are confident that significant fluid accumulation in the abdomen was unlikely to occur in the time interval between paracentesis and measurements.

**Pulmonary volumes**

A redistribution of lung volumes occurred: IC increased, whereas ERV and FRC decreased. In conditions such as ascites or pregnancy, the decrease in FRC is thought to be determined by an increased intra-abdominal pressure, which elevates the diaphragm in the thorax [1, 19]. Consistent with this hypothesis are the observations [1–3] that after LVP the FRC and ERV increase, which is confirmed by the present study.

**Breathing pattern and arterial blood gas values**

Chronic hyperventilation with a resulting compensated respiratory alkalosis is a common observation in patients with cirrhosis [20–22]. The present data confirm this observation, the increase in \( V^E \) being due to an increase both of tidal volume and respiratory frequency. Several mechanisms have been proposed to explain hyperventilation in cirrhotic patients [20, 21], but the real reason is still obscure. No change in \( V^E \) was observed after LVP, and this suggests that ascites was probably not implicated in this abnormality. Hypoxaemia is also frequently associated with cirrhosis and different mechanisms have been thought to be involved in
determining it: ventilation/perfusion (V’A/Q’) mismatching; intrapulmonary shunt; limitation of oxygen diffusion; and inadequate vascular tone [22, 23]. Two patients exhibited mild hypoxaemia (Nos. 4 and 8), but on average $P_{a,0_2}$ was normal and unchanged after LVP.

**Inspiratory muscle strength**

The patients studied showed normal or decreased inspiratory muscle strength, which did not change with LVP. These findings are consistent with those of HOURANI et al. [24], who measured respiratory muscle strength in 116 patients with severe liver disease of varying aetiology and found a decreased maximal inspiratory pressure (MIP) in 56% of them. The reasons as to why inspiratory muscle strength may be decreased in cirrhotic ascitic patients are not clear. In fact, the effects of increased abdominal pressure on diaphragmatic strength are controversial. In patients undergoing continuous ambulatory peritoneal dialysis (CAPD), a situation that from a mechanical point of view is similar to ascites, PREZANT et al. [25] reported that an increase of peritoneal dialysate from 0 to 3 L induced progressive decrease of FRC and increase both of MIP and $P_{du}$, measured in sitting position. Two factors have been thought to be involved in determining the increase in diaphragmatic strength in patients with raised intra-abdominal pressure [25, 26]: 1) the elevation of the diaphragm in the thorax by the high intra-abdominal pressure could lengthen diaphragmatic fibres, increasing the area of apposition of the diaphragm to the rib cage and improving the length-tension relationship of the diaphragm; and 2) the presence of liquid in the abdomen could decrease the abdominal compliance, thus providing a more effective fulcrum for the diaphragmatic action.

Based on the above considerations, one could expect that diaphragmatic strength is increased in ascitic patients and decreases with LVP. However, in the patients studied, LVP induced inconsistent changes. At variance with PREZANT et al. [25], SIAFARAS et al. [27] found that MIP, measured in sitting position, was low during CAPD and increased after drainage of the fluid. There are several reasons that could explain these discrepancies. Firstly, the positive effect of diaphragmatic elevation may be counterbalanced by an enlargement of the lower rib cage, thus, overly compliant [4]. Consistent with this hypothesis, the observation that a low $P_{ga,sw}$ was accompanied on inspiratory muscle force probably depends on the balance between the above-mentioned factors, and this may explain the different results observed in the present study and in other studies [25, 27].

The lack of effects of LVP on inspiratory muscle strength suggests that other mechanisms linked to liver disease are probably involved in determining inspiratory muscle weakness. High sericomicotonic diaphragmatic alterations characteristic of liver cirrhosis (such as hypoalbuminaemia and electrolyte abnormalities) might play a role, as confirmed by the observation of muscle wasting in patients with liver diseases [24], and, in particular, in cirrhotic patients [30]. Finally, one has also to consider that $P_{du, max}$ is an effort-dependent manoeuvre, so that variations in effort pre- and postparacentesis might influence the measure. Due to the clinical condition of our patients, we could not perform phrenic nerve stimulation, which is the only way to obtain more reliable measures of diaphragmatic strength.

**Chest wall mechanics**

Our patients showed high $P_{pl,sw}$ and relatively low $P_{ga,sw}$, both of which decreased significantly with LVP. Figure 3 shows that an end-inspiration, pleural pressure was normal or only slightly altered, while at end-expiration it was markedly increased. This increase appears to be the consequence of the high intra-abdominal pressure, as clearly indicated by the close relationship between the amount of fluid removed and decrease both in $P_{pl,EE}$ and $P_{pl,sw}$ (fig. 5a and b). Swings of $P_{du}$ were also high, indicating an increased activation of inspiratory muscles, and decreased significantly with LVP (figs. 2d and 3), this reduction being mainly due to a decreased $P_{du, EI}$ (fig. 3). Consistent with the observation of a significant reduction both in lung elastic load and threshold load after paracentesis, a decrease in overloading and, thereby, a reduced activation of inspiratory muscles are likely explanations for the decrease of $P_{du, EI}$ and $P_{du,sw}$ after LVP. The absence of significant modifications in Vahl/P$_{ga}$ and previous observations in experimental animals [29] argue against the possibility that liquid subtraction induced an increase of abdominal compliance, causing a lesser impediment to inspiratory diaphragmatic descent and, thus, resulting in a greater diaphragmatic shortening and the development of a lower $P_{du, EI}$ and $P_{du,sw}$. Finally, the finding of a high $P_{du,sw}$ accompanied both by high $P_{pl,sw}$ and low $P_{ga,sw}$ suggests an important contribution of the rib cage inspiratory muscles to generation of inspiratory pressure.

The magnetometry data show that the generation of tidal volume was contributed mainly by the abdomen and the relative contributions of abdomen and rib cage did not change with LVP. This is consistent with the findings of HANSON et al. [4], who observed in ascitic patients a relationship between lung volumes and intra-abdominal pressure, but, surprisingly, no change in intra-abdominal pressure with paracentesis. The explanation for this finding was that in ascitic patients abdominal muscles within the chronically distended abdominal wall are stretched beyond their elastic limit and are, thus, overly compliant [4]. Consistent with this hypothesis, the observation that a low $P_{ga,sw}$ was accompanied
by a high abdominal contribution to tidal volume genera-

tion confirms that abdominal compliance is probably

very high in these patients. Finally, the lack of changes

both in the contribution of the abdomen to the genera-

tion of tidal breathing and in the $V_{ab}/P_g$ with LVP

argues against change in abdominal compliance with

LVP, and supports the hypothesis of HANSON et al. [4].

Lung mechanics

Another important finding of this study is the ele-

vated load on the inspiratory muscles due both to an in-

creased pulmonary dynamic elastic load ($E_{dyn}$) and the

presence of a threshold load (PEEPi). In order to over-

come the increased lung mechanical load, ascitic pa-

tients have to increase the $P_{plaw}$ at each breath (table 3).

The interrelationships between changes in $P_{plaw}/V_T$ and

changes in $E_{dyn}$ and PEEPi indicate that after para-

centesis the load on the inspiratory muscles decreased and

that a lower $P_{pl}$ was necessary to generate the same

$V_T$. The following two factors are likely to contribute

to the increased $E_{dyn}$ in cirrhotic patients. Firstly, clo-

sure of alveolar units; this mechanism is thought to be

involved in causing a decrease in lung compliance both

in obese patients [31, 32], and in normal subjects dur-

ing chest wall strapping [33, 34]. The closure of air-

ways in cirrhotic patients may be a consequence of the

high intra-abdominal pressure and the consequent in-

creased pleural pressure (see below). Secondly, increase

in lung elastic recoil due to interstitial pulmonary oede-

ma, which is thought to be present in patients with liver

cirrhosis [35].

An interesting finding was the presence of a PEEPi.

Although the reasons for the presence of a PEEPi in

cirrhotic patients with ascites are complex, one has to

consider many possibilities. Firstly, an apparent PEEPi

may be due to a discrepancy between pleural and oeso-

phageal pressure in the supine position, in which the

present measurements were carried out. In fact, in this

position oesophageal pressure may be higher than pleu-

ral pressure [36, 37]. However, this cannot explain the

presence of a time lag between the beginning of the ne-

gative deflection in pleural pressure and the beginning

of inspiratory flow. Secondly, contraction of abdominal

muscles during expiration may result in an apparent

PEEPi or contribute to it [17]. This mechanism did not

appear to contribute substantially to the production of

PEEPi in the present patients, as they did not show an

increase in gastric pressure on expiration (fig. 1).

Thirdly, high intra-abdominal pressure causes pleural

pressure to be high at end-expiration (fig. 4), and this

in turn can determine an early closure of the airways.

As a consequence, the alveoli of the dependent lung

regions do not empty during expiration and a positive

pressure remains in them at end-expiration. This hypo-

thesis is supported by the close interrelationship be-

tween changes in PEEPi, the amount of fluid removed

and $P_{pl,EE}$ (figs. 5a, 5c and 6): the larger the amount

of fluid removed, the greater the reduction in $P_{pl,EE}$ and

PEEPi, and the lower the $P_{pl,EE}$, the lower the PEEPi.

Thus, we think that increased abdominal pressure play-

ed a major role in determining the presence of PEEPi.

However, other mechanisms, such as mechanical com-

pression of the small airways by dilated blood vessels

and interstitial oedema, cannot be ruled out [35].

Finally, the observation that LVP had small effects on

respiratory mechanics when less than 5–6 L of ascites

was removed (fig. 5a–c), suggests that a substantial

reduction of respiratory load can be achieved only if

large volumes of ascites are subtracted. This observa-

tion may have importance in the clinical management of

ascitic patients.

In conclusion, in supine cirrhotic patients, tense asci-

tes determines an overload for inspiratory muscles due

both to a high lung elastic load and the presence of

positive end-expiratory alveolar pressure. Large volume

paracentesis unloads the inspiratory muscles and de-

creases their activation, thus significantly improving the

respiratory function. Decreased inspiratory muscle strength,

possibly determined by the metabolic alterations char-

acteristic of liver cirrhosis, may be present in some

patients in the sitting position and is not modified by

large volume paracentesis.

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