Hypoxaemia and cirrhosis of the liver

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Funahashi, A., Kutty, A. V. P., and Prater, Sandra L. (1976). Thorax, 31, 303–308. Hypoxaemia and cirrhosis of the liver. In order to determine the frequency of hypoxaemia and to evaluate the role of increased closing capacity in producing hypoxaemia in patients with cirrhosis of the liver, 13 patients with well-established cirrhosis were studied. Arterial blood gases, spirometry, lung volume, and closing capacity measurements were made with the patients in the seated and recumbent positions after exclusion of cardiopulmonary dysfunction. Four of 13 and six of 12 patients exhibited significant hypoxaemia in the seated and recumbent positions respectively. Five of 13 patients showed a closing capacity greater than predicted. This frequency of increased closing capacity was not higher than in a group of smokers of the same age. Unlike Ruff et al. (1971), we did not find a consistent relationship between hypoxaemia and closing capacity.

The association between hypoxaemia and cirrhosis of the liver has been known since the report of Snell in 1935. Abnormal oxygen dissociation was initially thought to be the cause of this phenomenon, but subsequent studies showed that the change was not sufficient to explain the degree of hypoxaemia (Heinemann, Emirgil, and Mijnssen, 1960; Caldwell, Fritts, and Cournand, 1965). Portopulmonary or intrapulmonary shunts were postulated as a primary cause of hypoxaemia in cirrhotic patients since anatomical communications between the portal vein and the pulmonary venous system were observed in necropsy material (Calabresi and Abelmann, 1957) and produced in experimental animals (Khaliq, Kay, and Heath, 1972). The physiological significance of this shunt flow was studied by Nakamura et al. (1965) who found no positive correlation between such flow and hypoxaemia. The existence of intrapulmonary shunts has been well documented (Rydell and Hoffbauer, 1956; Hansoti and Shah, 1966; El Gamal et al., 1970), but its incidence was found to be very low (Berthelot et al., 1966; Stanley, Ackrill, and Wood, 1972).

In recent years ventilation-perfusion abnormality has been considered to be the major mechanism in causing hypoxaemia among patients with cirrhosis of the liver (Karetzky and Mithoefer, 1967; Cotes et al., 1968).

Since the introduction of the measurement of closing volume, it has been suggested that a smaller functional residual capacity (FRC) than closing capacity (CC) may be a cause of ventilation-perfusion abnormality leading to hypoxaemia (Craig et al., 1971) on the basis that if CC is greater than FRC the airways of dependent parts of the lung are closed during a part of the normal respiratory cycle. If perfusion continues through these areas during airway closure, it will result in a shunt and lead to arterial hypoxaemia. The CC tends to remain unchanged for both sitting and recumbent positions while FRC usually is smaller in the recumbent than in the sitting position. Therefore, the CC may exceed FRC only in the recumbent position (Leblanc, Ruff, and Milic-Emili, 1970). This relationship has been suggested as a mechanism for hypoxaemia seen only in the recumbent position (Hamosh and DaSilva, 1973).

Ruff and his associates (1971) studied the closing capacity of cirrhotic patients who had hypoxaemia and found that the CC was uniformly increased. They further reported that the CC was greater than FRC in most of their patients.

The purpose of the present study was to determine the frequency of hypoxaemia in patients

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Thorax (1976), 31, 303.
with cirrhosis of the liver who had no appreciable cardiopulmonary abnormalities, and to determine the significance of the FRC–CC value in producing hypoxaemia.

**MATERIAL AND METHODS**

Thirteen patients were selected for the study. All had cirrhosis of the liver, 10 had moderate to marked ascites, and three had no ascites. The criteria for selection were: (1) cirrhosis of the liver verified by biopsy; (2) absence of cardiopulmonary abnormalities including airways obstruction; and (3) normal chest radiograph. Informed consent was obtained from all subjects.

Spirometry was measured with a Stead-Wells spirometer and the lung volumes were determined by the helium dilution method (Meneely et al., 1960). The closing volume was measured by the single breath nitrogen washout method described by Buist and Ross (1973) using an expiratory flow rate below 0·3 l/s. The closing volume was that between the point of terminal rise in nitrogen concentration and the end of expiration. Closing volume plus residual volume was termed closing capacity. Only tracings showing a distinct rise in N2 concentration were used, and tracings which had a vital capacity deviating more than ±10% from that of the initial measurement with the Stead-Wells spirometer were discarded. Closing capacity is expressed either as a per cent of total lung capacity or as an actual volume. The coefficient of variation of repeated measurements of closing capacity/TLC in normal subjects in our laboratory, 8·7%, is similar to that reported by others (McFadden, Holmes, and Kiker, 1975).

Arterial blood was obtained from the brachial artery by an arterial puncture with a 20 gauge needle, and oxygen tension (Pao2), carbon dioxide tension (Paco2), and pH were measured with the Radiometer gas analyzer. All measurements were performed in both sitting and recumbent positions. The sequence of study was: (1) collection of arterial blood, (2) spirometry, (3) lung volume measurements, and (4) closing volume. The patients assumed each position for a minimum of 15 minutes before the start of the study to allow the redistribution of perfusion.

Eleven healthy smokers of age comparable to that of the cirrhotic patients were selected to measure the effect of cigarette smoking alone on closing volume and arterial blood gases as well as to see the relationship between FRC and CC. The predicted values for spirometry were based on the Veterans Administration and Army Cooperative Study (Kory et al., 1961).

**RESULTS**

Table I shows the age, smoking history, and spirometric measurements of 13 cirrhotic patients. Cases 3, 4, and 10 showed moderate reduction of FVC, but FVC was normal or near normal in the remaining cases. In all cases forced expiratory volume in one second (FEV1)/FVC was greater than 70%, indicating that there was no large airway obstruction. Table II shows the age, smoking history, spirometric measurements, and Pao2 of 11 healthy smokers. FVC and FEV1/FVC were normal except in cases 2 and 9, who had a minimally low FEV1 and FEV1/FVC ratio, respectively. Arterial oxygen tension (Pao2) was generally above 80 mmHg except for cases 2 and 9, who had slight hypoxaemia.

The Pao2 for both sitting and recumbent positions are shown in Figure 1. The shaded area

<table>
<thead>
<tr>
<th>TABLE I</th>
<th>AGE, SMOKING HISTORY, AND SPIROMETRY OF PATIENTS WITH CIRRHOSIS OF THE LIVER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case</td>
<td>Age</td>
</tr>
<tr>
<td>Cirrhotic patients with ascites</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>56</td>
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<tr>
<td>2</td>
<td>52</td>
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<tr>
<td>3</td>
<td>50</td>
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<tr>
<td>4</td>
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<td>8</td>
<td>68</td>
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<tr>
<td>9</td>
<td>66</td>
</tr>
<tr>
<td>10</td>
<td>48</td>
</tr>
</tbody>
</table>

| Cirrhotic patients without ascites | | | | |
| 11 | 48 | 20 | 4·59 (87) | 79 |
| 12 | 51 | 30 | 4·01 (105) | 81 |
| 13 | 57 | 30 | 3·05 (94) | 85 |
| Mean | 52 | 27 | 83 | |

1Pack years.
2Percent of predicted value.
### Table II

<table>
<thead>
<tr>
<th>Case</th>
<th>Age</th>
<th>Smoking</th>
<th>FVC</th>
<th>FEV1/FVC</th>
<th>PaO2</th>
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<td>30</td>
<td>4-10 (105)*</td>
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<td>93</td>
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<tr>
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<td>56</td>
<td>3-76 (82)</td>
<td>76</td>
<td>75</td>
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<tr>
<td>3</td>
<td>44</td>
<td>28</td>
<td>5-78 (116)</td>
<td>75</td>
<td>93</td>
</tr>
<tr>
<td>4</td>
<td>54</td>
<td>76</td>
<td>3-61 (94)</td>
<td>80</td>
<td>111</td>
</tr>
<tr>
<td>5</td>
<td>53</td>
<td>35</td>
<td>4-80 (111)</td>
<td>84</td>
<td>86</td>
</tr>
<tr>
<td>6</td>
<td>59</td>
<td>10</td>
<td>3-76 (94)</td>
<td>81</td>
<td>116</td>
</tr>
<tr>
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<td>40</td>
<td>36</td>
<td>4-30 (91)</td>
<td>80</td>
<td>86</td>
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<tr>
<td>8</td>
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<td>100</td>
<td>3-90 (90)</td>
<td>84</td>
<td>85</td>
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<tr>
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<td>34</td>
<td>4-57 (112)</td>
<td>69</td>
<td>76</td>
</tr>
<tr>
<td>10</td>
<td>59</td>
<td>21</td>
<td>3-90 (93)</td>
<td>77</td>
<td>84</td>
</tr>
<tr>
<td>11</td>
<td>50</td>
<td>12</td>
<td>4-35 (98)</td>
<td>78</td>
<td>91</td>
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<tr>
<td>Mean</td>
<td>52</td>
<td>40</td>
<td>99%</td>
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<td></td>
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</table>

*Pack years.

*Percent of predicted value.

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**FIG. 1.** Level of arterial oxygen tension of cirrhotic patients in both sitting and recumbent positions. An arrow indicates the change from sitting to recumbent position; — indicates no change: ○ patient with ascites and normal PaO2; ● patient with ascites and hypoxaemia; △ patient without ascites and normal PaO2; ▲ patient without ascites and hypoxaemia.

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is mean ±1 SD for PaO2 in healthy subjects (Sorbini et al., 1968). In the sitting position four of 10 patients with ascites showed hypoxaemia and three of them had PaO2 values below 66 mmHg. None of the patients without ascites showed hypoxaemia. Most of the patients with ascites showed a slight hyperventilation, with arterial carbon dioxide tension (Paco2) in the range of 31 to 35 mmHg, except case 9, who showed a marked hyperventilation (Paco2 of 23 mmHg and PaO2 of 120 mmHg). There was no appreciable change in Paco2 between the sitting and recumbent position in any patient studied. In the recumbent position for which the changes of PaO2 from the sitting position are indicated by the arrow in Fig. 1, five of nine patients with ascites showed hypoxaemia. One patient (10), who was hypoxaemic in the sitting position, was unable to assume the recumbent position because of a large amount of ascites and, therefore, was not included. Three patients, two with ascites and one without ascites, became hypoxaemic in the recumbent position. The degree of hypoxaemia was not altered significantly with a change in body position for three subjects who were hypoxaemic while both sitting and recumbent. Figure 2 shows the CC of all 13 patients, superimposed on the shaded area which indicates the predicted normal value according to Buist (1973). Five of 10 ascitic patients showed an increased CC, whereas all three non-ascitic patients were within the normal range. Of the five who had increased CC, two had hypoxaemia, whereas two patients with hypoxaemia had a normal CC. Of 11 healthy smokers measured for this study, five showed an increased CC. In two normal smokers with hypoxaemia, one had an increased CC while the other had a normal CC.

The relationship between FRC and CC in cirrhotic patients is shown in Figure 3. Most of the patients in the sitting position who had a normal PaO2 had FRC greater than CC. In the four patients who had a low PaO2, however, two showed FRC greater than CC (FRC–CC >0), and two showed FRC smaller than CC (FRC–CC <0). In the recumbent position most of the hypoxaemic patients had an FRC value smaller than CC (FRC–CC <0). However, there was a wide scatter...
of the FRC-CC value in patients who showed normal \( P_{\text{ao2}} \) in the recumbent position. Figure 4 shows this FRC-CC value among the healthy smokers. Again there was a wide scatter of this value. Two subjects who had hypoxaemia showed FRC value smaller than CC. On the other hand, four subjects without hypoxaemia had FRC-CC value below zero.

**FIG. 4.** FRC-CC value (litre) of healthy smokers in sitting position: ○ subject with normal \( P_{\text{ao2}} \); ● subject with hypoxaemia.

**FIG. 3.** FRC-CC value (litre) of cirrhotic patients in both sitting and recumbent positions. An arrow indicates the change from sitting to recumbent: ○ patient with ascites and normal \( P_{\text{ao2}} \); ● patient with ascites and hypoxaemia; △ patient without ascites and normal \( P_{\text{ao2}} \); ▲ patient without ascites and hypoxaemia.

**DISCUSSION**

The incidence of hypoxaemia among patients with cirrhosis of the liver has been reported in various series (Blackburn *et al.*, 1960; Rodman, Sobel, and Close, 1960; Bashour, Miller, and Chapman, 1961), though pulmonary disease or congestive heart failure, both of which may cause hypoxaemia, were not carefully excluded. In the present study we excluded hypoxaemia due to cardiopulmonary dysfunction, yet still found the frequency of hypoxaemia to be surprisingly high. Since many patients with cirrhosis have associated pulmonary disease or congestive heart failure, the overall prevalence of hypoxaemia would be even higher.

Hypoxaemia was more frequent in the recumbent position. The degree of hypoxaemia, however, did not differ in two patients with ascites who were hypoxaemic in both sitting and recumbent positions, and one patient who was hypoxaemic only in the recumbent position did not have ascites. These observations suggest that the mechanical disturbance of diaphragmatic movement due to ascites was probably not a major factor in producing hypoxaemia among these patients. It has been well established that smokers have a higher incidence of increased CV and CC (Buist, 1973; Funahashi, Melville, and Hamilton, 1975). In the present series, the incidence of increased CC among the subjects with cirrhosis of the liver was no greater than that in healthy smokers with a comparable smoking history. This suggests that cirrhosis of the liver, even with the presence of a moderate to marked degree of ascites, did not specifically alter the CC. That conclusion differs from that of Ruff *et al.* (1971), though the reason is not apparent. It may be due to the difference of subjects studied or the different method used to measure the closing capacity. In their series, none had ascites and all but two had hypoxaemia, whereas a majority of our patients had moderate to marked ascites and only four had hypoxaemia in a sitting position. Two of these four hypoxaemic patients, however, had normal CC. If high CC were secondary to increased interstitial fluid around the small airways, as postulated by Ruff *et al.* (1971), one might expect to see any effect of cirrhosis exaggerated by the presence of ascites.

Another aspect of the present study that needs to be considered is the possible effect of lung restriction due to ascites upon calculation of CC. The CC is calculated by adding RV to the closing volume and dividing by TLC. Therefore, if TLC decreased while RV remained unchanged, a net effect of low TLC upon calculation of CC would be an erroneous increase of calculated CC. In our series, RV was essentially within the normal range in those cases where there was a significant reduction of TLC. Since the frequency of increased
CC in our cirrhotic patients was no greater than that in healthy smokers, this possible effect of restrictive change due to ascites on the calculation of CC appears to be minimal.

Ruff et al. (1971) used a bolus gas technique whereas we used a single breath nitrogen washout method. These two methods are based on different principles but correlate well in healthy subjects (Travis, Green, and Don, 1973). It is, however, not certain that they correlate in disease. The possibility that a change of regional RV caused by ascites could have resulted in a low CC measurement is not entirely ruled out.

We were unable to find a correlation between FRC and CC relationships and hypoxaemia. This led us to search our files for this relationship among healthy smokers. To our surprise, four smokers out of 11 examined had a normal PaO₂ even though their FRC was much smaller than CC. Only two subjects with FRC smaller than CC exhibited hypoxaemia. It is, therefore, evident that having a CC larger than FRC does not necessarily result in hypoxaemia, although hypoxaemic subjects who were cirrhotic or were smokers tended to have FRC values smaller than CC. This lack of correlation may reflect the fact that lowered alveolar PaO₂ causes vasoconstriction and results in a redistribution of perfusion. Hypoxaemia may result only when, for some reason, this compensatory mechanism is lost and the under-ventilated areas continue to be perfused. It has been suggested that in patients with cirrhosis of the liver this regulatory mechanism may be lost (Daoud, Reeves, and Schaefer, 1972).

The present study suggests that the cause of hypoxaemia among cirrhotic patients is not directly attributed to an increased closing capacity. Severe hypoxaemia observed occasionally by others was probably due to a pulmonary capillary shunt, while in a large percentage of patients with slight to moderate hypoxaemia the most likely cause is a failure to control perfusion locally, leading to an increased ventilation-perfusion disproportion. In some patients this ventilation-perfusion mismatching may be the result of an increase in CC relative to the FRC.

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REFERENCES


Requests for reprints to: Dr. A. Funahashi, Pulmonary Function Laboratory, Veterans Administration Center, Wood, Wis. 53193, USA.
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