Lung sound intensity in patients with emphysema and in normal subjects at standardised airflows

H J W Schreur, P J Sterk, J Vanderschoot, H C J van Klink, E van Vollenhoven, J H Dijkman

Abstract
Background A common auscultatory finding in pulmonary emphysema is a reduction of lung sounds. This might be due to a reduction in the generation of sounds due to the accompanying airflow limitation or to poor transmission of sounds due to destruction of parenchyma. Lung sound intensity was investigated in normal and emphysematous subjects in relation to airflow.

Methods Eight normal men (45–63 years, FEV₁, 79–126% predicted) and nine men with severe emphysema (50–70 years, FEV₁, 14–63% predicted) participated in the study. Emphysema was diagnosed according to pulmonary history, results of lung function tests, and radiographic criteria. All subjects underwent phonopneumography during standardised breathing manoeuvres between 0·5 and 2 l below total lung capacity with inspiratory and expiratory target airflows of 2 and 1 l/s respectively during 50 seconds. The synchronous measurements included airflow at the mouth and lung volume changes, and lung sounds at four locations on the right chest wall. For each microphone airflow dependent power spectra were computed by using fast Fourier transformation. Lung sound intensity was expressed as log power (in dB) at 200 Hz at inspiratory flow rates of 1 and 2 l/s and at an expiratory flow rate of 1 l/s.

Results Lung sound intensity was well repeatable on two separate days, the intraclass correlation coefficient ranging from 0·77 to 0·94 between the four microphones. The intensity was strongly influenced by microphone location and airflow. There was, however, no significant difference in lung sound intensity at any flow rate between the normal and the emphysema group.

Conclusion Airflow standardised lung sound intensity does not differ between normal and emphysematous subjects. This suggests that the auscultatory finding of diminished breath sounds during the regular physical examination in patients with emphysema is due predominantly to airflow limitation.

Pulmonary emphysema is associated with morphological lesions within the acini of the lung.¹ It is defined as a condition characterised by abnormal permanent enlargement of the airspaces distal to the terminal bronchioles, accompanied by destruction of their walls, without obvious fibrosis.² The diagnosis of emphysema during life is based on clinical history, physical examination, chest radiography, and lung function testing.³ A traditional characteristic feature of emphysema is the auscultatory finding of diminished intensity of lung sounds.⁴ This can be caused either by poor transmission of sounds as a result of parenchymal destruction or by reduced generation of sounds due to airflow limitation.⁵

Phonopneumography has shown that even normal subjects have considerable intersubject and intrasubject variability in the intensity of the inspiratory vesicular sounds heard on the chest wall.⁶ In patients with emphysema this variability seems to be much greater.⁷ Ploysongsang et al found that lung sound transmission is often abnormal in patients with emphysema, being reduced in some areas of the lung but normal or even increased in other areas.⁷ In addition, they observed that regional breath sounds vary from breath to breath.⁷ When measured at various locations on the chest, regional sound intensity appeared to be related to regional ventilation,⁸ which points to a potential role of airflow limitation in the reduction of lung sounds in emphysema.

In healthy human volunteers lung sound intensity is highly dependent on airflow at the mouth.⁹ The frequency spectrum of lung sounds, however, does not seem to be affected by airflow.¹⁰ These findings were extended by other studies, indicating that lung sound intensity increased with the square of both inspiratory and expiratory flow.¹¹,¹² Airflow limitation therefore might be one of the major determinants of diminished breath sounds in patients with emphysema.

The objective of the present study was to test the hypothesis that lung sound intensity is similar in normal subjects and in patients with emphysema when measured at equal airflow rates. We therefore measured lung sound intensity by airflow standardised phonopneumography in normal and emphysematous men. As the within subject variability of lung sound intensity has been reported to be relatively high,¹³ we also determined the
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Table 1 Characteristics of the subjects

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<th>TLCO (% pred)</th>
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FEV₁-pre—forced expiratory volume in one second, measured before bronchodilatation; TLCO—transfer factor for carbon monoxide; N—normal; E—emphysema; a—a antitrypsin deficiency; +—non-smoker; ——smoker; ex—ex-smoker; IA—inhaled anticholinergic; I₂E—one inhaled β₂ adrenergic; OX—oral xanthine derivative; Crom—sodium cromoglycate.

Methods

DESIGN OF THE STUDY

The study consisted of two parts.

Part 1 For the comparison of lung sound intensity in normal and in emphysematous subjects we had a screening day (for checking the inclusion criteria) and a study day for both groups. On the study day phonopneumography was carried out while subjects were breathing at standardised flow rates.

Part 2 The repeatability of the measurements was investigated in normal subjects, who visited the laboratory on two separate days one to three days apart. On each day they underwent airflow standardised phonopneumography.

SUBJECTS AND INVESTIGATIONS

Part 1 Eight normal subjects (45–63 years) and nine patients with severe emphysema (50–70 years) participated in this study. The normal subjects were recruited from hospital personnel and acquaintances and the patients were selected from outpatients of the department of pulmonology. The normal subjects had no history of lung disease, no abnormalities found by physical examination, a normal chest radiograph, and normal forced expiratory volume in one second (FEV₁: 79–126% predicted) (table 1). The patients with emphysema had had clinical symptoms of emphysema, such as regular wheezing and prominent dyspnoea after exercise, for several years. Physical examination showed tachypnoea, prolonged expiration, a hyperresonant chest, and diminished breath sounds on regular auscultation. The diagnosis of emphysema was based both on the appearance of the chest radiograph (paucity of peripheral arteries and abnormal length and width of the lungs and size of the retrosternal space, heart size, and diaphragm position) and on the results of lung function tests. All patients had a decreased FEV₁ (14–63% predicted) and the measurements of FEV₁ after inhalation of 200 μg salbutamol showed that little of the decrease was reversible—less than 10% of the predicted value (table 1). In addition, transfer factor for carbon monoxide (TLCO), measured by the single breath holding technique, was reduced in all patients (21–68% predicted). The clinical condition of the patients was stable, and none had had symptoms of respiratory tract infections during the two weeks before the tests. At the time of the study treatment was continued as usual (table 1). The study was approved by the hospital ethics committee, and all subjects gave informed consent.

Part 2 The repeatability of the measurements of lung sound intensity was examined in 10 normal male volunteers (23–42 years, FEV₁ 91–124% predicted) with no history of pulmonary disease. The subjects were recruited from hospital personnel. All were nonsmokers, and at the time of the study none of them used any medication.

PHONOPNEUMOGRAPHY

The experiments included synchronous recordings of airflow and lung volume changes at the mouth, obtained by spirometry (Morgan Spiroflow, UK), and phonopneumography in a sound proof room. Lung sounds were recorded with four identical air coupled piezoelectric microphones (Sony ECM-150T) at standardised locations on the right chest. Microphone 1 (Mic₁) and microphone 2 (Mic₂) were attached over respectively the 2nd and the 5th intercostal space at the midclavicular line, microphone 3 (Mic₃) over the 4th intercostal space at the midaxillary line, and microphone 4 (Mic₄) over the 9th intercostal space at the midscapular line. The air coupled microphones (sensitivity 2.5 mv/Pa signal to noise ratio >40 dB (0.1 Pa, 1 kHz), band width 300 Hz –20 kHz ± 3 dB) were mounted in stainless steel housings that were fixed to the chest wall.
with adhesive electrocardiography rings. The combinations of microphones and housings had a flat frequency response from 150 Hz to 3 kHz (±3 dB). The output signals from the four microphones were passed through identical amplifiers and fourth order Bessel high pass filters with a cut off frequency of 337.5 Hz to obtain very effective removal of heart sounds and motion noises. A headphone connection was provided for monitoring lung sounds from two channels at a time. Airflow, lung volume changes, and the signals from the four microphones were sampled at 5000 Hz each, and stored on the hard disk of an IBM PC-AT computer.

To standardise the breathing manoeuvres flow-volume loops were displayed in front of the subjects on an oscilloscope screen (Hewlett-Packard HP 1741A). First the subjects were asked to inhale towards total lung capacity (TLC), which was used as a reference volume. Subsequently they performed breathing manoeuvres between TLC-0.5 l and TLC-2 l with inspiratory and expiratory target flows of 2 and 1 l/s respectively in cycles of 3 seconds during 50 seconds.

ANALYSIS
For each microphone airflow dependent power spectra were computed by means of the fast Fourier transform method. The power spectra were analysed on 100 ms lung sound intervals, a Hanning window being used. These intervals were centred around lung sound samples that were corresponding in time to airflow samples at which the airflow was a multiple integer of 0.1 l/s. The spectra obtained in this way were averaged between all complete breathing cycles of one registration for each distinct airflow value, and for the ascending and the descending limb separately. This resulted in three dimensional diagrams of an averaged breathing cycle, showing the relation between airflow (in l/s, x axis), lung sound frequency (in Hz, y axis), and the logarithm of the lung sound intensity (in dB, z axis), and separately for the rising and the descending limb of the flow curve (fig 1). Fast Fourier transform spectra were determined for every 0.1 l/s. To express lung sound intensity within and between the groups of subjects, the log power at 200 Hz was measured at inspiratory airflows of 2 l/s (LSI20i) and 1 l/s (LSI10i) and at an expiratory airflow of 1 l/s (LSI10e). At 0 l/s the slope of the airflow versus time curve is rather steep. Thus the interval of 100 ms centred on this airflow will comprise lung sounds generated at airflow values from about 0.35 l/s on inspiration to 0.55 l/s on expiration. Unfortunately, this is inevitable when fast Fourier transforms are used if frequency resolution is not to be lost. For this reason 0 l/s has not been used for the statistical comparison.

The frequency of 200 Hz was chosen as the maximal energy of lung sounds has been reported to occur from 116 to 350 Hz13-17 whereas muscle and heart sounds have frequencies predominantly below 100 Hz.18 We decided to measure the intensity at one frequency, as the power spectra in the three dimensional plots from all subjects, both normal and emphysematous, were very similar in shape, and very smooth. Furthermore, during inspection of the three dimensional diagrams and the multidimensional wave forms there was no evidence for crackles20 or wheezes21 that could influence the intensity of the lung sounds.

The repeatability of the results of lung sound intensity measurements was computed by using 95% confidence intervals of the differences between day 1 and day 222 and by using the intraclass correlation coefficient obtained by analysis of variance for repeated measurements.23 The intraclass correlation reflects the ratio of the between subject variability to the between subject plus within subject variability of the measurements. The differences in lung sound intensity between normal subjects and patients with emphysema were analysed by multivariate analysis of variance, with airflow, microphone and group as independent variables. We considered p values less than 0.05
Figure 2  Mean lung sound intensity with standard errors in eight normal and nine emphysematous subjects for each of the microphone locations (Mic,) and three levels of airflow (1 l/s expiratory, 1 l/s inspiratory, and 2 l/s inspiratory).

Results

PART 1

Satisfactory recordings could be obtained in all subjects. Representative examples of three dimensional diagrams in one normal (top) and one emphysematous subject (bottom) are shown in figure 1. The frequency spectra of the lung sounds appeared to be similar in normal and emphysematous subjects and ranged from 40 to 1100 Hz, the highest frequencies occurring during inspiration at 2 l/s.

There was a considerable variability in the measured lung sound intensity both within and between subjects. Lung sound intensity was significantly influenced by microphone location (p < 0.005) and by airflow (p < 0.005). In both groups of subjects lung sounds from the microphone placed midclavicularly over the second intercostal space (Mic,) were the loudest, whereas lung sounds from the microphone placed under the armpit (Mic,) were the weakest (fig 2). Lung sound intensity was greatest at 2 l/s inspiratory flow, less at 1 l/s inspiratory flow, and least at 1 l/s expiratory flow (fig 2). When these effects of microphone location and airflow were taken into account, however, there was no significant difference in lung sound intensity between the normal and the emphysema group (p = 0.72; fig 2).

PART 2

When the repeatability of the recordings was analysed there was no significant difference in lung sound intensity between the phonopneumographic registrations on the two days (p = 0.61). The identity plot of the results of lung sound intensity measurements between day 1 and day 2 is shown in figure 3. The mean of the differences (with 95% confidence interval, CI) between repeated measures of intensity was found to be -0.34 (95% CI 5.37) dB. The intraclass correlation coefficient varied between microphone locations and airflow levels (table 2). It ranged from 0.49 for Mic, at 1 l/s expiratory flow to 0.82 for Mic, 1 l/s inspiratory flow, with two outliers—0.16 for Mic, at 2 l/s inspiratory flow and 0.36 for Mic, at 1 l/s expiratory flow. The relatively low intraclass correlations for these measurements appear to be due to a very limited between subject
variability in conjunction with the usually observed within subject variability (fig 3).

Discussion
The results of the present study indicate that airflow standardised measurements of lung sound intensity are feasible and reproducible. Sound intensity is dependent on airflow and varies between different locations on the chest. For a given level of airflow, however, there is no difference in lung sound intensity between normal subjects and patients with emphysema at any of the investigated locations. This suggests that the auscultatory finding of diminished lung sounds in emphysema is predominantly due to concurrent airflow limitation. The reduction of the intensity of lung sounds in patients with emphysema has been extensively reported. In these publications, however, breathing manoeuvres were not standardised, nor was sound intensity measured at specific levels of airflow. With the traditional approach there appeared to be an inverse relation between lung sound intensity and the degree of airflow limitation in patients with airways obstruction. Thus it is not entirely unexpected that sound intensity in patients with emphysema is similar to that in normal subjects when recorded at the same level of airflow.

The present results might be affected by methodological errors—for example, in selection of subjects, methods of measurements, or analysis. Firstly, the diagnosis of emphysema was based on pulmonary history and findings from the physical examination, lung function values, and the chest radiograph. It could be argued that emphysema is a morphological diagnosis that cannot be made without pathological evidence from tissue samples. The specificity of the chest radiograph for the pathological lesions is, however, sufficiently large to justify its use in confirming the presence of moderate to severe emphysema. The present selection criteria do not allow us to specify the subtype of emphysema any further.

Secondly, the microphones, their localisation, and signal processing were standardised as much as possible. Each microphone was consistently used at the same standardised location. Lung sound intensity was determined at 200 Hz, which represents the centre of the range of frequencies observed by others to contain the maximum energy of lung sounds. In addition, despite high pass filtering at 337.5 Hz the maximum energy in our three dimensional diagrams coincides with about 200 Hz. The spectra obtained from the subjects were very similar in shape and (as a result of the averaging of the spectra) were very smooth, so that choosing only a single frequency for the determination of lung sound intensity will hardly influence the accuracy of the measurements. Further, inspection of the three dimensional diagrams enabled us to exclude the occurrence of adventitious lung sounds that could be responsible for energy peaks at other frequencies. We therefore assumed the measured power at 200 Hz to be representative of the overall sound intensity at the specific flow rate.

High pass filtering of lung sounds at 337.5 Hz to obtain a very effective elimination of heart and muscle noise did not invalidate measurements of lung sound intensity at 200 Hz in the present study. When a high pass filter is used, by definition signals at the cut off frequency are attenuated by 3 dB. Above this frequency the attenuation decreases to 0 dB, and below this frequency the signals are increasingly attenuated. When a 337.5 Hz fourth order Bessel high pass filter is used, lung sounds at 200 Hz were attenuated to the same extent in all subjects (9-6 dB). Thus the ratio of lung sound intensity observed in normal subjects to that in patients with emphysema has not been affected. As we measured intensity as log power, the differences of lung sound intensity (in dB) between subjects will not have changed either.

Thirdly, the present data on the repeatability of lung sound intensity measurements confirm the validity of our methods. The intraclass correlation showed that the between subject variability in lung sound intensity is sufficiently large in relation to the total variability (between and within subjects), even within this relatively homogeneous group of normal subjects. Further, the power analysis showed that a difference in lung sound intensity of at least 7.16 dB could have been detected between the normal and the emphysematous subjects in the present study. This difference is sufficiently small in relation to the effects of the other determinants of lung sound intensity, such as location on the chest wall and airflow (fig 2). The absence of diminished lung sounds in emphysema in the present study is therefore unlikely to have been caused by methodological errors.

How can we explain the similarity of lung sound intensity in normal and emphysematous subjects? Normal "vesicular" breath sounds are considered to originate predominantly from complex turbulence within the central airways. Minor contributions to normal breath sounds may be generated by unsteady movement of vortices formed at junctions in the fifth to the 13th generations in the human bronchial tree. It has, however, been suggested that inspiratory vesicular lung sounds are partly generated by other still unexplained mechanisms. We do not know whether any of these mechanisms is influenced by the mechanical changes within the lung during the development of pulmonary emphysema. The disease is characterised by parenchymal destruction leading to alveolar enlargement and loss of alveolar attachments to the bron-
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chiroles.18 This is reflected physiologically by a decrease in lung elastic recoil pressure, an increase in lung volume, and airflow limitation.1 Thus the parenchymal destruction may change the transmission of lung sounds, whereas the abnormalities in the airways may affect their generation.

Firstly, the parenchymal tissue serves as an important conducting medium, along which sounds can propagate towards the chest.17 18 Thus the alveolar destruction and enlargement in emphysema potentially compromises lung sound transmission. It has been observed that lung sounds may diminish with increasing lung volume.15 16 19 Even though we standardised the target volume in our experiments from 0-5 to 2 l below TLC, the absolute lung volume in the patients might still have been greater than in the normal subjects. Nevertheless, we found no difference in lung sound intensity between the two groups, which indicates that radio graphically confirmed parenchymal destruction did not alter lung sound intensity in our patients with emphysema. This is in accordance with the observation that the transmission of artificial sounds introduced at the mouth may be either decreased or increased in emphysema.9 Reduced sound transmission therefore does not seem to have a major influence on the auscultatory findings in these patients.

Secondly, any changes in airway geometry may alter the generation of sounds. In emphysema the static shape of the intrapulmonary airways is irregular and tortuous.30 This might enhance sound production during inspiration and expiration.14 17 18 On the other hand, the airways may be obstructed in emphysema, as a result of dynamic compression or mucus thickening or both, caused by the disease process itself or by concomitant bronchitis or bronchiolitis.30 Apart from potentially causing adventitious lung sounds, this obstruction results in airflow limitation and reduced ventilation, which has been observed to diminish lung sound intensity.15 16 17 Indeed, Ploysongsang et al16 showed that regional lung sound intensity is correlated with regional ventilation in emphysema, which suggested the hypothesis that an airflow dependent reduction in sound generation could explain the auscultatory findings in this disease. Our present observations favour this hypothesis. Lung sound intensity was dependent on airflow, but appeared to be normal in emphysema when airflow was strictly standardised. Apparently, all other remaining mechanical abnormalities in emphysema do not substantially contribute to the intensity of lung sounds on the chest.

The results of the present study have clinical implications. The reduction in lung sound intensity has been considered as a major clinical criterion for the diagnosis of pulmonary emphysema.2 4 This auscultatory finding, however, appears to be highly dependent on airflow, which is likely to be less in patients with emphysema than in normal subjects during the usual physical examination procedures. When airflow is standardised the abnormality disappears, making any inference based on auscultatory examination of lung sound intensity in the clinical diagnosis of emphysema questionable.

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