Clinical application of regional lung function studies in infants and small children using $^{13}$N

R. RONCHETTI*, JANET STOCKS, NANETTE FREEDMAN, H. GLASS†, and S. GODFREY

*From the Department of Paediatrics and Neonatal Medicine, and Department of Medical Physics, Hammersmith Hospital, London

Ronchetti, R., Stocks, J., Freedman, N., Glass, H., and Godfrey, S. (1975.) Archives of Disease in Childhood, 50, 595. **Clinical application of regional lung function studies in infants and small children using $^{13}$N.** A technique is described for the investigation of regional lung function in infants and children using $^{13}$N and a gamma camera. Boluses of isotopic gas are inhaled and perfused while the lung fields are scanned. The child is lightly sedated and breathes normally throughout. Regional function is assessed in terms of the distribution of gas and blood, and the balance between ventilation and perfusion is estimated by comparing an index of the ventilation per unit volume of ventilated lung with that of perfused lung. The use of the method in 8 infants and children with different clinical problems is described to show its application. The method is capable of defining the severity and localization of any abnormality and may also be useful in showing normal function in suspect areas.

Respiratory disease is a major cause of morbidity and mortality during infancy and early childhood. Great advances have been made over the years in developing methods for measuring lung function in infants analogous to those used in adults, and normal ranges have been defined (Godfrey, 1974). These tests are only practicable during the first few months of life when the infant can be lightly sedated and studied without active co-operation. Above this age the infant or small child is very liable to resist any attempts to measure breathing and such studies are virtually impossible without anaesthesia until about 5 years of age. Unfortunately, a number of conditions present or progress in this period of 6 months to 5 years when children are least accessible. Moreover, conventional lung function tests or blood gas measurements can only assess total lung function, and give no clues as to regional variations in function which may be clinically important.

In adults it is now routine practice to assess regional lung function after inhalation and infusion of radioactive gases such as xenon ($^{133}$Xe). Originally such measurements were made with several external counters (Knipping et al., 1955), but more recently a gamma camera has been used (Secker-Walker et al., 1973). Some preliminary work has been carried out in healthy and sick infants (Ronchetti et al., 1971; Koch et al., 1973), and recently a more definitive method of using $^{133}$Xe in children has been described giving some normal data (Treves et al., 1974). We decided to develop the use of radioactive nitrogen ($^{15}$N), which has a half-life of 10 minutes, because of its lower solubility in tissue and fat compared with xenon which renders it more suitable for studying lung function in infants and children with pulmonary problems. The method has been designed so that active co-operation is not needed. We present the results of our initial studies in 8 infants and children where the method proved to be of considerable clinical value.

**Subjects and methods**

The 8 children, who were all undergoing investigation or treatment were between 2 weeks and 2-2 years old; one child was studied twice during his illness. Some clinical details of the patients are given in Table I. A variety of problems was studied, each baby being investigated to help establish the diagnosis or severity of
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TABLE I

Details of patients

<table>
<thead>
<tr>
<th>Case no.</th>
<th>Diagnosis</th>
<th>Age</th>
<th>Weight (kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Agenesis of the R lung</td>
<td>2 w</td>
<td>2·1</td>
</tr>
<tr>
<td>2</td>
<td>Cystic lesions of lung with generalized</td>
<td>(a)</td>
<td>5 w</td>
</tr>
<tr>
<td></td>
<td>hyperinflation and Fallot's tetralogy</td>
<td>(b)</td>
<td>5·6</td>
</tr>
<tr>
<td>3</td>
<td>Pulmonary hyperinflation, airway</td>
<td>26 m</td>
<td>11·0</td>
</tr>
<tr>
<td>4</td>
<td>Cystic fibrosis</td>
<td>8 w</td>
<td>3·6</td>
</tr>
<tr>
<td>5</td>
<td>Atypical arthrogryposis with kyphoscoliosis</td>
<td>24 m</td>
<td>7·5</td>
</tr>
<tr>
<td>6</td>
<td>Possible congenital lobar emphysema with</td>
<td>9 w</td>
<td>3·5</td>
</tr>
<tr>
<td></td>
<td>collapse of R upper lobe</td>
<td>11 w</td>
<td>5·6</td>
</tr>
<tr>
<td>7</td>
<td>Alveolar hypoplasia</td>
<td>5 m</td>
<td>2·6</td>
</tr>
<tr>
<td>8</td>
<td>Wilson-Mikity syndrome</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The conditions listed in Table I. $^{13}$N was produced by the Medical Research Council Cyclotron Unit and piped to a dispensing laboratory adjacent to the room where the gamma camera was situated. The desired quantity of radioactivity was made up as a gas or dissolved in sterile isotonic saline ($0·1-0·5$ mCi/ml) immediately before use. During and after inhalation or intravenous administration of the radioisotope, the lungs were examined continuously with the gamma camera for 5 minutes, and the data stored on digital magnetic tape and subsequently processed by computer (Fig. 1). The data were corrected for physical decay of the radioisotope and for the natural background count-rate. The count-rates at selected time intervals were then shown on an oscilloscope and presented graphically. The computer program allowed considerable automation of the data processing. Of particular advantage was the availability of a light pen and oscilloscope which allowed selected regions of any shape to be delineated within the lung fields and the appropriate count-rate versus time curves to be obtained easily for each selected region of the lung.

The smaller infants were lightly sedated with chloral hydrate ($30$ mg/kg) and studied after a feed, while the older babies usually received diazepam as premedication. The patient lay supine over the gamma camera and was made comfortable so that he dozed off or lay quietly. The position of the infant was checked with a radioisotope marker in all but the earliest studies. In order to inject $^{13}$N intravenously, a scalp vein needle was inserted in the smaller infants and a needle was placed in an arm vein in the older infants. A continuous infusion of dextrose-saline was used to keep the line patent until the injection was given. For the inhalation study we initially used a hood placed over the head through which a stream of test gas was blown. In later studies we found it more satisfactory to inject the gas through a nasal catheter passed back as far as the nasopharynx. The babies tolerated both the intravenous and the nasal catheter well and breathed spontaneously throughout.

**Fig. 1.—Diagram illustrating general principles of technique of $^{13}$N studies.**
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During the studies, expired radioactive gas was expelled to the outside air by means of a simple pump and wide-bore tube placed close to the baby's face.

For each infant $^{13}$N was given once by inhalation and once by infusion with a suitable time period between each procedure to allow all the residual radioactivity to clear. In the initial studies an attempt was made to deliver the gas for each test at a constant rate for one minute as suggested by Ronchetti et al. (1974), but it was apparent from the records that this gave an inadequate count-rate for subsequent data analysis. The highest count-rates were observed after the catheters were eventually flushed through with air or saline, effectively delivering a bolus of gas. This procedure was therefore abandoned and a bolus of gas was administered in the six later studies. The inhalation bolus (4 ml gas) was delivered into the nasopharynx at the end of an expiration and the infusion bolus was injected as rapidly as possible (4 ml $^{13}$N 'dissolved' in saline delivered over approximately 2-3 s). The dose of radioactivity was calculated on the basis of 0.3 mCi/kg entering the lungs for each inhalation or perfusion study up to a maximum of 2 mCi. The total calculated radiation received by the lungs for a combined inhalation and perfusion study was 180-200 mrad for infants and approximately 100-130 mrad for older children. The maximum tissue doses allowing for the worst case of a 50% right-left shunt were <25 mrad. This compares favourably with other isotope studies used in children (Seltzer, Keriakes, and Saenger, 1964), and with diagnostic radiology such as bronchography which results in about 3 times as much irradiation. No infants appeared to suffer any ill-effects from the study.

**Results**

The individual values for the calculated indices for each baby are given in Table II. As each patient presented a different problem, each study will be briefly described.

**Case 1.** Was one of monozygotic twins and clinically and radiologically appeared to have agenesis of the right lung (Fig. 2). A $^{13}$N scan confirmed the absence of any perfused lung tissue on the right and an even distribution of blood to the left (and only) lung. The ventilation of this perfused lung tissue was normal (see below). The inhalation study was technically unsatisfactory. 14 weeks after the investigation the infant died a cot death and necropsy confirmed pulmonary agenesis.

**Case 2.** The second infant to be studied will be described in detail elsewhere (Godfrey et al., 1975). He had Fallot's tetralogy of a mild degree but radiological evidence of cystic lesions, especially at the lung bases (Fig. 3). The possibility of a localized congenital abnormality was considered and a $^{15}$N scan was carried out at the age of 5 weeks. Both inhalation and perfusion studies had poor count rates but showed relatively even distribution indices. There was extremely poor ventilation of perfused lung compared to ventilated lung, confirming generalized abnormality with gas trapping. Subsequently this infant was restudied at the age of 6½ months using the improved bolus technique. The results for the ventilation of perfused lung were similar and the difference between ventilation of ventilated lung and the ventilation of perfused lung was even more striking and can be seen in the shape of the respective washout curves (Fig. 4). Though the upper zones were relatively less affected than the other zones (Table II) the
Case 3. This infant was referred for study because of unexplained pulmonary hyperinflation and fixed airways obstruction, thought to be related to previous bronchiolitis. The study showed a marked discrepancy between the right and left lungs, with nearly twice as much gas and blood going to the right. Ventilation of ventilated and perfused lung was also better on the right but poor in all areas, especially for perfused lung. These findings suggested the presence of Macleod's syndrome (Macleod, 1954) which had not previously been suspected. The abnormality predominantly affected the right lung but there was generalized damage to both lungs as a result, presumably, of the bronchiolitis.

Case 4. This infant had meconium ileus due to cystic fibrosis but had not yet had any radiological evidence of lung damage at the time of the 13N scan. The inhalation study was poor because of bad positioning and only one region on the left could be defined. Both inhalation and perfusion studies appeared to be normal except that there was some inequality in the regional distribution of perfusion. The study confirmed the absence of significant pulmonary involvement.

Case 5. This male was 2 years old at the time of study and had atypical arthrogryposis with kyphoscoliosis making regional localization difficult. It was thought that his right lung was under-perfused. The 13N study showed an uneven pattern in which the only definite abnormality was poor ventilation of perfused lung at the right base. Underperfusion of the right lung could not be substantiated.

Case 6. A female was born at 34 weeks' gestation and had the respiratory distress syndrome from
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**Fig. 2.**—X-rays of monozygotic twins. Twin on the right, Case 1, had agenesis of the right lung and was studied by the $^{13}$N technique.

**Fig. 3.**—Case 2. Chest x-ray of infant with hyperinflated lungs and radiolucent areas especially at the bases, together with Fallot's tetralogy.
which she recovered after $O_2$ therapy. At 5 weeks of age she had an acute respiratory illness, possibly due to aspiration, and needed artificial ventilation for 48 hours. She remained tachypnoeic and chest x-rays showed persistent overinflation of the left upper lobe and collapse of the right upper lobe (Fig. 5). The possibility of congenital lobar emphysema was considered and the question of an exploratory thoracotomy was discussed. She was referred for a preliminary $^{13}$N investigation. This gave unequivocal evidence of reduced distribution of gas and blood to the right upper zone and entirely normal function of the overinflated left upper lobe (Table II). Clearly, this was incompatible with compression of the right lung by an emphysematous left lung, and suggested that the collapse on the right was the primary with compensatory overdistension of the normal left lung. The baby was treated by vigorous physiotherapy and the right upper lobe re-expanded with concomitant return of the left lung to a normal size.

**Case 7.** A baby of 11 weeks with hypothyroidism and hypoxia of uncertain aetiology. The $^{13}$N scan showed relatively even distribution of gas and blood, but a markedly reduced ventilation of perfused lung compared with ventilated lung in all areas (Table II). This was interpreted as evidence for very poor ventilation-perfusion matching which would be expected with interstitial lung disease. The excellent washout of inhaled $^{13}$N seemed to preclude any airway disease. A subsequent lung biopsy showed generalized pulmonary hypoplasia. (A repeat $^{13}$N study after treatment with thyroxine, not included in the present series, showed significant improvement.)

**Case 8.** A preterm infant with mild respiratory
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are given and they do not describe how they obtained the necessary co-operation. We elected to make measurements during spontaneous respiration in order to have a simpler and practical test able to furnish high counts with small doses of radioactivity. We initially tried to achieve steady state conditions with relatively short inhalations and perfusions, but this method did not give satisfactory count-rates with the dose of radioactivity which we were prepared to use at the time. It could well be argued that steady state inhalation would provide a better index of the function of ventilated lung than a bolus inhalation and might give a better measure of lung volume than the area of the oscilloscope image. Based on our present experience we hope to explore a suitable steady state inhalation procedure. When using bolus injections (Heck-scher et al., 1966) the shape of the input function of the bolus arriving at the region of the lung which is being examined is not certain and some radioactivity is removed by ventilation while the latter part of the bolus is still arriving. Though exact analysis of the bolus infusion data is complicated, nevertheless the amount of radioactivity lost by ventilation during arrival of the bolus over a few seconds is probably very small and will hardly influence the results.

The mathematical quantitation of the results in the present study is subject to the limitations mentioned above, but for clinical purposes it seemed adequate to treat all the data in this way. Exponential analysis of washout curves (Fowler, Cornish, and Kety, 1952) has been used in many radioisotope studies of lung function, but this requires the subdivision of the lungs into arbitrary 'slow' and 'fast' compartments, and may be associated with significant errors (Glass and de Garreta, 1971). The fractional ventilation index which we used has the advantage of providing an indication of the total function of each region in terms of the average regional ventilation per unit volume. Within the limits discussed we believe this to be an acceptable approach. The distribution indices based on peak heights are also an approximation since they depend upon the volume of lung in each region. The use of the area of the oscilloscope picture of the region is a relatively poor substitute, but not unreasonable when it is realized that the thinner upper portion of the upper zone is more or less equivalent to the thinner lower portion of the lower zone around the dome of the diaphragm. In fact the results which we got in normal regions of lung seem to justify the method, at least for clinical purposes, and they resemble the distribution indices corrected for volume used by

Discussion

We believe that the $^{15}$N technique was useful in the diagnosis and evaluation of function in the 8 infants described.

Radioisotopic evaluation of lung function in adults is an established procedure. They are normally studied after rebreathing in order to equilibrate all areas (Secker-Walker et al., 1973) and to correct the results for the volume of lung being viewed (Ball et al., 1962). A similar approach has been used in children (Treves et al., 1974), but their technique must require 2 or 3 minutes of rebreathing through a face-mask though no details

Fig. 5.—Case 5. Chest x-ray of infant with hyperinflation of the left upper lobe apparently causing mediastinal deviation and compression of right lung.

distress syndrome at birth who went on to develop clinical and radiological features suggestive of the Wilson-Mikity syndrome though the referring hospital questioned whether the changes could have been secondary to localized infection. Regular lung function tests had shown hyperinflation and increased airways resistance. As in the previous patient, the $^{15}$N investigation showed a relatively even distribution of gas and blood but with ventilation-perfusion imbalance shown by the very poor ventilation of perfused lung. Unlike the previous baby, the ventilation of ventilated lung was also relatively poor, suggesting generalized lung and airway disease. The widespread nature of the changes would not be compatible with local damage from infection and strongly supported the diagnosis of the Wilson-Mikity syndrome.
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Treves et al. (1974). The same regions should be used for both the ventilation and the perfusion studies if possible so that comparisons can be made. It is possible that the use of an area overestimates volume in diseased regions where the overall size of the lung is retained (e.g. bronchiectasis), but this becomes a semantic problem in the sense that our index then gives the distribution of blood per unit 'volume' of tissue, which will be low, rather than per unit volume of gas-filled lung. Clearly both types of result could well be of use, and we hope to explore this in the future. The fractional ventilation is of course entirely independent of the volume measurement.

The distribution index for inhaled gas is related to the fractional ventilation of ventilated lung because both must reflect regional ventilation. However, the distribution index after injection reflects pulmonary blood flow, while the fractional ventilation of perfused lung is related to the balance between ventilation and perfusion. When gas trapping is present, the ventilation study usually results in a smaller visible area than the perfusion study, and errors would occur if this smaller area were used, because it would exclude some lung tissue which was perfused but poorly ventilated.

We did not consider it acceptable to carry out studies in healthy infants and so we do not have any normal values. However, because of the localized nature of the disease in some of the infants, it has been possible to calculate the fractional ventilation for apparently healthy regions of ventilated and perfused lungs. The value of the mean ventilation per unit volume index for 5 'healthy' ventilated regions was 4·27 (range 3·36–5·05), and the fractional ventilation index for 'healthy' perfused regions was 3·09 (range 2·08–3·90). These fractional ventilations can be compared with the values for ventilated lung of 1·95 found in 20 adults by Secker-Walker et al. (1973) and the values for perfused lung of 1·68–2·00 found in 11 normal adults by Heckscher et al. (1966), using different methods. Theoretically, the fractional ventilation should resemble alveolar ventilation divided by end-expiratory lung volume. Using the mean data related to age collected by Godfrey (1974), this index is calculated to be 4·00 for the newborn infant, 3·85 for children between 1 and 5 years, and 2·25 for adults. These theoretical values are similar to those we found in children and others found in adults, and thus serve as some justification for the methods.

The distribution indices which we found in healthy regions were usually close to the ideal value of 1·0 (e.g. left lower zone in Case 6, Table II) and this agrees with the results in normal lungs found by Treves et al. (1974). Their fractional ventilation indices appear to have been normalized for whole lung clearance and cannot be compared with our values or with the likely values based on alveolar ventilation and lung volume.

The value of the 13N studies in the individual patients is largely self-evident from the results, and will not be discussed in detail. It seems that the test is most useful in defining the extent of a known abnormality (e.g. Case 2) or showing the localization of an abnormality (e.g. Cases 3, 5, and 6), but can also help in differential diagnosis (Cases 7 and 8) and even in showing healthy lung tissue (Cases 1 and 4). Though the bulk distribution of gas and blood were useful indices in some cases (3 and 6), the demonstration of gas trapping and hence poor ventilation-perfusion relations was often the most helpful finding (e.g. Cases 2, 7, and 8). This emphasizes the need for both inhalation and perfusion studies.

Finally, we recommend the use of 13N, which is a more physiological radiisotope tracer than 133Xe because of its very low solubility. Its safety of handling due to its short half-life is an additional advantage. The major disadvantage is of course that it can only be used in an institute where there is a cyclotron on site. The number of medical cyclotrons around the world has now reached 14 (Glass, 1973). Our technique could be adapted to use 133Xe, but the computation of results is likely to be more difficult because of tissue solubility. We have subsequently studied a further 21 infants and children using the bolus technique and have confirmed the clinical value of the method as illustrated in these first 8 patients.

We are grateful to our colleagues in the Medical Research Council Cyclotron Unit for help in performing these studies, and especially to Mr. P. Buckingham. We also thank Drs. E. N. Hey, E. Shinebourne, and D. Lawson for referring their patients to us for investigation. We have benefited from the advice of a number of colleagues and would particularly like to thank Drs. J. M. B. Hughes, J. B. West, and P. Winlove.

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Clinical application of regional lung function studies using $^{13}$N


Correspondence to Dr. S. Godfrey, Department of Paediatrics and Neonatal Medicine, Hammersmith Hospital, Du Cane Road, London W12 0HS.
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