Disturbance in respiratory mechanics in infants with bronchiolitis

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ABSTRACT The passive flow-volume and partial forced expiratory flow-volume techniques were used to assess pulmonary function in 14 spontaneously breathing infants with acute respiratory syncytial virus bronchiolitis. Two additional infants were studied while paralysed and ventilated. During the acute stage of the illness there was a significant reduction in forced expiratory flow rates and an increase in respiratory resistance. Although the mean thoracic gas volume for the group was increased, five infants did not compensate for their airways obstruction by hyperinflation. Curvilinear passive flow-volume curves were seen in three of the 14 non-ventilated infants and in both ventilated infants. At follow up three to four months later all passive flow-volume curves were linear. There was a significant reduction in hyperinflation and an increase in forced expiratory flow rates, but values still differed significantly from those in normal infants.

Introduction

Acute viral bronchiolitis is the most common serious lower respiratory infection in the first six months of life. The clinical picture is one of small airway obstruction and pulmonary hyperinflation. Pulmonary function tests in the acute phase of the illness confirm the clinical and radiological evidence of airflow obstruction and gas trapping. Criticism has, however, been levelled at the tests used to assess lung function, suggesting that they may not reflect changes in airway function accurately.

New methods of evaluating lung function in young infants have been developed recently. Total respiratory system compliance and resistance have been measured by the occlusion and passive flow-volume techniques in newborn and older infants and maximum expiratory flow rates by the partial forced expiratory flow-volume technique. The two techniques allow measurements to be made without oesophageal balloons, which have major methodological problems in infancy, and without the complex equipment needed for the rebreathing method for measuring airways resistance, which is not particularly suitable for ill infants. The passive technique makes use of the observation that infants have a Hering-Breuer reflex. After transient occlusion at end inspiration the infant expires passively; the normal plot of expiratory flow against lung volume is a straight line. This allows calculation of the resistance and compliance of the respiratory system without use of invasive techniques.

In the present study we combined the two techniques to study infants with acute bronchiolitis due to respiratory syncytial virus during the acute and recovery phases of the illness. In addition, passive expiration was studied in two paralysed and ventilated infants with bronchiolitis. The aim was to document abnormalities in respiratory mechanics and to obtain insights into mechanisms used by the infants to compensate for the pathological changes.

Methods

PATIENTS We studied 16 previously healthy infants admitted to hospital with acute bronchiolitis. The project was approved by the hospital ethics committee and informed consent obtained from the parents. In most cases one parent was present during the examination.

The mean age of the 14 infants studied at about the eighth day of the illness was 20 (range 4–41) weeks. They were studied again three to four months later. Two infants aged 6 and 8 weeks were studied while paralysed and ventilated. The diagnosis of bronchiolitis was based on the presence of tachypnoea,
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hyperinflation, wheezing, and widespread crepitations. These were present at the time of the first study, though the spontaneously breathing infants no longer required nursing in a high oxygen environment. All patients had a positive result in the immunofluorescence test for respiratory syncytial virus antigen in the nasopharyngeal secretions. In the follow up study eight babies still had intermittent wheeze but were otherwise well.

INVESTIGATIONS

The spontaneously breathing infants were studied lying supine with the neck slightly extended in a 40 litre body plethysmograph, after receiving 80 mg/kg chloral hydrate. A three position slide valve, modified from that originally described by Le Souef et al, was fixed to the face with silicone putty. Flow was measured with a Fleisch No I pneumotachograph and a Validyne DP 45 pressure transducer. Volume was determined by electronic integration of the flow signal and drift was carefully adjusted. Flow and volume were displayed on a Tektronix 5223 digitising oscilloscope and recorded on tape with a Racial thermionic store 4D recorder. These signals and pressure were also displayed on a chart recorder (HP 7754A). Mouth pressure was measured via a port between the face mask and slide valve by means of a Hewlett Packard 128 OC pressure transducer. The pressure in the plethysmograph and cuff used for compression were measured with similar transducers.

Respiratory timing

During spontaneous tidal ventilation the duration of a respiratory cycle (Ttot) and inspiratory time (Ti) were measured from the flow tracings and expressed as Ti/Ttot.

Passive expiratory flow manoeuvre

The passive compliance (Crs), resistance (Rrs), and time constant (Trs) of the total respiratory system were obtained by the airway occlusion technique. Brief occlusion at end inspiration was performed with the slide valve between mask and pneumotachograph. The occlusion was released as soon as a plateau of mouth pressure was reached and, in the absence of flow, equalisation of pressure within the respiratory system was assumed. The linear part of the passive expiratory flow-volume (PEFV) curve was extrapolated to the flow and volume axes to calculate Crs, Rrs, and Trs (fig 1). (Compliance (Crs) = extrapolated expired volume (Ve)/occlusion mouth pressure (Pao), and resistance (Rrs) = occlusion mouth pressure (Pao)/extrapolated expiratory flow (Vo) at end inspired volume.)

In the two paralysed and ventilated infants the slide valve was attached to the endotracheal tube and ventilating bag, which bypassed the pneumotachograph. Constant pressure tracings during iso-volume ventilation and pressure plateau during occlusion excluded any appreciable leak around the endotracheal tube or in the measuring device. Because of the prolonged expiration to zero flow in these infants with severe airways obstruction, oxygen enriched air was supplied before each measurement and skin colour and heart rate were constantly monitored. Occlusion was performed at several different lung volumes, but before each occlusion the lungs were inflated to total lung capacity to standardise preceding pressure-volume relationships. PEFV curves were recorded after removal of the occlusion and the expiration was allowed to continue until there had been no expiratory flow for 3 to 4 seconds. The PEFV curves obtained in an individual patient initiated from different lung volumes were compared by assuming that after each passive expiration the same absolute end expiratory volume was achieved. We found that the PEFV curves of an individual infant were superimposed, except for the transient rise in flow immediately after occlusion. The transient flow recorded immediately after removal of the occlusion was ignored and Trs, Rrs, and Crs were calculated for each lung volume at which the occlusion was performed.

Forced expiratory flow manoeuvre

Partial forced expiratory flow-volume (FEFV) curves and maximum flow at functional residual capacity (VmaxFRC) were obtained by a modified version of the rapid compression technique. The chest and abdomen were compressed by rapid inflation of a plastic cuff enclosed in a nylon mesh jacket (fig 2). The cuff covered the chest and abdomen and its position did not change throughout the examination. The outer nylon jacket minimised pressure dissipation away from the chest. With this device about 70% of the cuff pressure was transmitted to the alveoli and airways as
measured by the increase in mouth pressure during occlusion.12 Virtually instantaneous inflation of the cuff was achieved by decompressing a gas storage drum through a wide bore connecting tube.12 Pressure within the drum was carefully regulated by a water filled manometer and a safety blow off valve. FEFV curves were obtained by progressively and carefully increasing chest compression until the shape of the FEFV curve and $V_{\text{maxFRC}}$ were constant and did not increase with further increases in chest compression pressure. Care was taken to avoid a negative "effort" effect when increasing pressure resulted in submaximum flow. With this occurred the cuff pressure was reduced until true maximum flow was again achieved. Final selection of curves with maximum flow for analysis was made when the raw signals were played back from the tape on to an oscilloscope.

Thoracic gas volume
Thoracic gas volume (TGV) was measured at end expiration in the constant volume body plethysmograph by the classic method of DuBois. As the infant attempted three or four spontaneous breaths after airways occlusion mouth pressure and box pressure were plotted on the $x$ and $y$ axes of a storage oscilloscope and TGV was obtained by applying Boyle's law. Calculations were corrected for dead space and adjusted for the difference between the measured and previously stable FRC. TGV at end inspiration was also measured in 10 babies during the first study, though unless otherwise stated TGV refers to end expiratory TGV.

Data analysis
Calculations were made after slowly playing back the taped records on to the oscilloscope and XY plotter (National VP 6123A). Each final measurement was a mean of at least five individual measurements made from technically satisfactory curves except in the case of $V_{\text{maxFRC}}$, where the highest value achieved was taken. The results were compared with normal data obtained by the same investigators from six healthy infants studied longitudinally on four occasions when aged 4-55 weeks. Prediction values based on height were derived from that study.8

The Wilcoxon signed rank and Mann-Whitney tests were used to compare the data from the infants with bronchiolitis and to compare these with the data from the normal infants. The 0.05 level of probability was assumed to be significant.

Results

Acute studies
Mean end expiratory TGV for the 14 infants was increased (table 1), though five infants had an end expiratory TGV within the normal range (less than 30 ml/kg). Mean (SEM) end inspiratory TGV minus tidal
volume in the 10 babies in whom this measurement was performed was 42 (3.5) ml/kg, significantly higher than end expiratory TGV (37 (3.5) ml/kg; p < 0.05). End inspiratory TGV less tidal volume was higher than end expiratory TGV in infants both with and without hyperinflation, though more strikingly in the former.

The passive expiratory flow-volume (PEFV) curve was a straight line in 11 of the 14 spontaneously breathing patients. In the remaining three patients it became curvilinear towards end expiration. Superimposing the forced, passive, and tidal volume curves produced similar lines for the three curves in these patients (fig 3).

Mean Rrs for the 14 infants was significantly increased, whereas mean specific compliance (sCrs) (that is, compliance divided by the lung volume at which it was measured) and mean specific conductance (sGrs) were significantly less than in the normal infants (table 1).

The FEFV curve was convex towards the x axis in 13 of the 14 patients. There was a significant decrease in the mean VmaxFRC and mean VmaxFRC/TGV (table 1). Twelve of the 14 infants had VmaxFRC/TGV below the 95% prediction interval (fig 4). The mean compression pressure needed to achieve Vmax in the patients was 35 cm H_2O compared with 34 cm H_2O in normal infants.

The five infants without hyperinflation had changes in VmaxFRC/TGV, mean sCrs, and mean sGrs similar to those of the infants with hyperinflation, though mean (SEM) Ti/Ttot was significantly shorter (0.382 (0.04) v 0.426 (0.03); p < 0.05).

In the two paralysed infants with severe bronchiolitis the PEFV curves were also curvilinear. Occlusion at different lung volumes allowed measurement of Rrs and Crs at varying lung volumes and showed increasing Rrs towards end expiration, followed by decreasing Crs at slightly lower lung volumes (fig 5).

**FOLLOW UP**

Lung function in most infants had improved when retested three to four months later. No PEFV curve remained curvilinear. TGV/kg was significantly less than during the acute phase of the illness (p < 0.05) and VmaxFRC/TGV was increased (p < 0.002). Trs and sGrs had not changed significantly, whereas sCrs had increased (p < 0.05). By comparison with a group of healthy infants (table 2), however, significant gas trapping and decreased volume corrected flow rates were still evident.

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**Table 1** Pulmonary function measurements (mean (SEM)) in 14 infants with acute viral bronchiolitis compared with measurements in six healthy infants

<table>
<thead>
<tr>
<th></th>
<th>Bronchiolitis</th>
<th>Normal infants</th>
<th>Significance of difference between groups (p)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>acute phase</td>
<td>(n = 14)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>(n = 6)</td>
<td></td>
</tr>
<tr>
<td>Age (w)</td>
<td>20-0 (3-4)</td>
<td>19-0 (1-8)</td>
<td>NS</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>6-22 (0-42)</td>
<td>6-90 (0-44)</td>
<td>NS</td>
</tr>
<tr>
<td>Height' (m')</td>
<td>0-23 (0-018)</td>
<td>0-27 (0-023)</td>
<td>NS</td>
</tr>
<tr>
<td>TGV (ml)</td>
<td>228 (19)</td>
<td>177 (11)</td>
<td>&lt; 0-05</td>
</tr>
<tr>
<td>TGV/kg (ml/kg)</td>
<td>37-6 (2-8)</td>
<td>25-8 (1-6)</td>
<td>&lt; 0-01</td>
</tr>
<tr>
<td>VmaxFRC (ml/s)</td>
<td>94 (15)</td>
<td>284 (28)</td>
<td>&lt; 0-001</td>
</tr>
<tr>
<td>VmaxFRC/TGV</td>
<td>0-406 (0-055)</td>
<td>1-594 (0-086)</td>
<td>&lt; 0-0001</td>
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<tr>
<td>Crs</td>
<td>6-84 (0-50)</td>
<td>9-04 (0-99)</td>
<td>NS</td>
</tr>
<tr>
<td>sCrs (cm H_2O^2)</td>
<td>0-031 (0-002)</td>
<td>0-051 (0-005)</td>
<td>&lt; 0-01</td>
</tr>
<tr>
<td>Rrs (cm H_2O.s.ml^-1)</td>
<td>0-049 (0-003)</td>
<td>0-039 (0-003)</td>
<td>&lt; 0-05</td>
</tr>
<tr>
<td>sGrs (s^-1.cm H_2O^-1)</td>
<td>0-099 (0-007)</td>
<td>0-150 (0-012)</td>
<td>&lt; 0-01</td>
</tr>
<tr>
<td>Trs (s^-1)</td>
<td>0-325 (0-021)</td>
<td>0-359 (0-056)</td>
<td>NS</td>
</tr>
</tbody>
</table>

For abbreviations see text.

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**Fig 3** Tidal, passive, and partial forced expiratory flow-volume curves from one infant. The passive curve was superimposed by extrapolating all curves to zero flow on the assumption that this occurs at the same absolute lung volume. Note the expiratory flow limitation in tidal and passive expiratory curves.
Discussion

This study has shown a pronounced reduction in forced expiratory flow and an increase in respiratory resistance in infants with viral bronchiolitis. Thoracic gas volume at functional residual capacity was substantially increased. Total respiratory compliance did not differ significantly from that of normal babies, though specific compliance was reduced, a reflection of the increase in TGV. The time constant of the respiratory system did not differ from that of normal infants. These findings are similar to those reported in studies using oesophageal balloons to measure intrathoracic pressure and the forced oscillation technique. The scientific validity of both these techniques has been questioned. The techniques used in this study have been developed over recent years. The forced expiratory technique was first applied to newborn infants and papers are now appearing on its use in older infants. One potential problem with this method is that the infant may start inspiration before reaching VmaxFRC. The consistency in the forced expiratory flow-volume curves in this study suggests that expiration concluded at a similar point with successive compressions. To obtain reproducible forced expiratory curves, it is important that the compression pressure is gradually increased until maximum flow is obtained, as was done in this study. The compression pressure needed to achieve maximum flow in the babies with bronchiolitis was similar to that in normal infants.

The original studies of the passive technique were also undertaken in newborn infants but there are now reports of its use in healthy older infants and in the evaluation of the efficacy of salbutamol in infants with acute viral bronchiolitis. It relies on the presence of the Hering-Breuer reflex and the assumption that after a period of short occlusion the infant passively expires to FRC. To exclude active respiratory muscle activity requires direct measurement of muscle activity, which is difficult in infants. There may be an initial flow

Table 2  Pulmonary function measurements (mean (SEM)) in 14 infants who had recovered from acute viral bronchiolitis compared with measurements in six healthy infants

<table>
<thead>
<tr>
<th></th>
<th>Bronchiolitis</th>
<th>Normal infants</th>
<th>Significance of difference between groups (p)</th>
</tr>
</thead>
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<tr>
<td></td>
<td>recovery phase</td>
<td>(n = 14)</td>
<td>(n = 6)</td>
</tr>
<tr>
<td>Age (w)</td>
<td>39.0 (3.6)</td>
<td>30.5 (0.34)</td>
<td>NS</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>8.95 (0.39)</td>
<td>8.55 (0.53)</td>
<td>NS</td>
</tr>
<tr>
<td>Height (m²)</td>
<td>0.37 (0.016)</td>
<td>0.36 (0.016)</td>
<td>NS</td>
</tr>
<tr>
<td>TGV (ml)</td>
<td>270 (19)</td>
<td>214 (16)</td>
<td>&lt; 0.05</td>
</tr>
<tr>
<td>TGV/kg (ml/kg)</td>
<td>30.3 (1.8)</td>
<td>25.0 (1.1)</td>
<td>&lt; 0.02</td>
</tr>
<tr>
<td>VmaxFRC (ml/s)</td>
<td>181 (27)</td>
<td>262 (30)</td>
<td>NS</td>
</tr>
<tr>
<td>VmaxFRC/TGV</td>
<td>0.675 (0.990)</td>
<td>1.238 (0.118)</td>
<td>&lt; 0.02</td>
</tr>
<tr>
<td>(TGV/s)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Crs (ml/cm H₂O)</td>
<td>10.24 (0.67)</td>
<td>10.28 (1.09)</td>
<td>NS</td>
</tr>
<tr>
<td>sCrs (cm H₂O²)</td>
<td>0.039 (0.003)</td>
<td>0.047 (0.002)</td>
<td>&lt; 0.05</td>
</tr>
<tr>
<td>Rs (cm H₂O.s.ml)</td>
<td>0.038 (0.003)</td>
<td>0.035 (0.003)</td>
<td>NS</td>
</tr>
<tr>
<td>sGs (s².cm H₂O²)</td>
<td>0.109 (0.011)</td>
<td>0.142 (0.015)</td>
<td>NS</td>
</tr>
<tr>
<td>Trs (s⁻)</td>
<td>0.388 (0.032)</td>
<td>0.354 (0.043)</td>
<td>NS</td>
</tr>
</tbody>
</table>

For abbreviations see text.
Disturbance in respiratory mechanics in infants with bronchiolitis

Thoracic gas volume was measured in the standard way used in this laboratory for 20 years. One group has recently reported low values of TGV in babies with clinically obvious hyperinflation and offered several possible explanations. Other laboratories, however, have confirmed hyperinflation in infants with airways obstruction. The suggestion that differences in calibration may explain the different findings is unlikely as similar methods have been used by the workers who found hyperinflation clinically and physiologically and by those who did not. Further work is necessary to account for these different findings.

Interestingly, end inspiratory TGV minus tidal volume was slightly greater than end expiratory TGV, which is the opposite of previous findings in infants with airways obstruction.

The most interesting findings in this study were the curvilinearity of the PEFV curves in three patients, the absence of hyperinflation in five, and the measurements made in the two ventilated paralysed infants. In normal infants the PEFV curve is a straight line. Linearity implies that the product of Crs and Rrs remains constant during passive expiration. The curvilinearity found in the patients with bronchiolitis indicates a changing time constant during expiration, probably due to varying speeds of emptying of areas of lung with different degrees of obstruction. In the paralysed infants there was a rapid rise in resistance with a fall in compliance towards smaller lung volumes, suggesting progressive narrowing and finally closure of small airways with air trapping. Glottic narrowing is suggested as another possible explanation but was excluded in our intubated, paralysed infants; so airways closure is the most likely explanation. Further, with glottic narrowing the initial expiratory resistance usually exceeds end expiratory resistance. Flattening of the PEFV curve therefore occurs at the beginning rather than towards end expiration. Another possible explanation for the curvilinearity is expiratory muscular activity during

 transient at the start of passive expiration, reflecting pressurisation of the pneumotachograph.

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![Fig 5 Passive expiration in one of the paralysed and ventilated infants: (a) curvilinear passive expiration (the bar below indicates the volumes of respirator tidal ventilation); (b) calculated Crs and Rrs at the different lung volumes during the passive expiration shown in (a).](image-url)
expiration.\textsuperscript{24} If this occurred early in expiration, an additional increase in mouth pressure should have been observed.

Eleven infants had linear PEFV curves in the presence of increased resistance and hyperinflation. The likely explanation is that tidal breathing was occurring at higher lung volumes where Rrs and Crs did not change during expiration. Possibly, however, increasing resistance was balanced by decreasing compliance during expiration so that their product remained constant.

The bar below the curvilinear PEFV curve for the paralysed and ventilated patient shown in figure 5 indicates the volume of ventilator tidal ventilation. It shows that an increase in end expiratory volume allowed breathing at a lower time constant. This leads to increased ventilation without any change in respiratory frequency. This compensation for airways closure may explain why no difference was found in time constants from the passive curve between infants with bronchiolitis and the normal controls. The forced expiratory flow-volume curve showed increased concavity with decreased VmaxFRC in the infants with bronchiolitis. This apparent discrepancy may have arisen because during passive expiration the increase in Rrs is balanced by a decrease in Crs, so that the time constant is unchanged. During the forced manoeuvre, however, the driving pressure for expiration is significantly decreased by the pressure transmitted from the cuff to the pleural space. In the presence of airways disease this pressure leads to narrowing or closure of airways (or both) with a resultant reduction in expiratory flow rates.

The reduction in VmaxFRC is most likely to be due to disease of intrapulmonary airways. It has been suggested that expiratory narrowing of the glottis, "laryngeal braking," may be important in maintaining hyperinflation in asthma,\textsuperscript{27} but if this was a factor in these babies some flattening of the initial portion of the PEFV curve and a much greater rise in Rrs would have been expected.

Five patients with flow limitation, two of whom had curvilinear PEFV curves, did not compensate by hyperinflation. This group appeared to maintain adequate ventilation by spending relatively more time on expiration. The reason for this mode of compensation was not clear. A similar pattern was seen in some patients recovering from acute asthma many years ago.\textsuperscript{28}

To analyse the Crs and Rrs when PEFV curves are curvilinear requires interruption during expiration, as in our paralysed infants. This analysis should also be possible in spontaneously breathing infants with the interruptor technique described for animals.\textsuperscript{28} With curvilinear PEFV curves calculation of Crs and Rrs from the initial part of passive expiration, which approximates to a straight line, may be misleading. Extrapolation of the straight part to the volume axis will underestimate the passive expired volume and a falsely low Crs will be calculated. Similarly, the calculated Rrs will underestimate the change in Rrs during expiration and so is valid only for the lung volume at end inspiration.

Thus our studies using recently developed techniques showed substantial reduction in forced expiratory flow in infants with acute viral bronchiolitis and suggest that this is due to changes in intrapulmonary airways. There was a much smaller change in Rrs, which is probably more influenced by disease in larger airways. Some infants did not compensate for airways narrowing with hyperinflation and they had severe expiratory flow limitation. Further investigation of why they failed to adopt this defence mechanism may give a better understanding of its origin. Although functional impairment was less at follow up three to four months after the acute bronchiolitis lung function was still significantly different from that of normal infants.

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References
11 Mansell AL, Bryan AC, Levison H. Relationship of lung
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Disturbance in respiratory mechanics in infants with bronchiolitis.
J Seidenberg, I B Masters, I Hudson, A Olinsky and P D Phelan

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