Cerebral blood flow velocity variability in infants receiving assisted ventilation

J M RENNIE, M SOUTH, AND C J MORLEY

University Department of Paediatrics, Addenbrooke's Hospital, Cambridge

SUMMARY Cerebral blood flow velocity was measured using Doppler ultrasonography in 20 babies who weighed less than 2500 g at birth and who were receiving assisted ventilation; their patterns of spontaneous respiration were recorded simultaneously. The babies were induced to breathe synchronously or asynchronously with the ventilator by manipulating the inspiratory and expiratory time settings. The variability of cerebral blood flow velocity (coefficient of variation) was calculated from the area of the maximum Doppler frequency shift envelope for 10 cardiac cycles from 211 recordings made on 42 occasions, and was greatest within 12 hours of birth after which it fell progressively over the next 48 hours.

Variability of cerebral blood flow velocity was significantly greater when the infants were breathing out of synchrony with the ventilator (median 11%, interquartile range 8–14%) than when they were either apnoeic (median 5%, 3–7%), or breathing synchronously with the ventilator (median 5%, 3–6%).

Perlman et al reported a large variability from beat to beat in the blood flow velocity in the cerebral arteries of some premature neonates using continuous wave Doppler ultrasonography. Twenty one of the 23 infants with wide variability developed intraventricular haemorrhages and possibly the transmission of fluctuations from the systemic arterial blood pressure to the capillaries of the germinal matrix led to their rupture. A pronounced reduction in the incidence of intraventricular haemorrhage was achieved by selective paralysis of infants with wide variability in blood flow velocity, with pancuronium. The authors suggested that the infants' respirations that occurred out of synchrony with the ventilator were responsible for this variability, and elimination of spontaneous respiration by pancuronium caused the reduction in the incidence of haemorrhage. Unfortunately no recordings of respiration were made to confirm this hypothesis. Pancuronium may have had its effect either by reducing non-respiratory movements or by some direct action on the circulation.

Selective paralysis with pancuronium of high risk infants may reduce the risk of intraventricular haemorrhage, but unfortunately the drug may have cardiovascular side effects and may necessitate the use of higher ventilator pressures to maintain adequate oxygenation. Alternative measures for infants who breathe against the ventilator are triggered ventilators or the induction of synchronous ventilation by manipulation of ventilator timing. Selective paralysis of infants who breathe out against ventilator inflations or who require defined peak pressures and rates has reduced the incidence of pneumothoraces but this has not been associated with a noticeable reduction in the incidence of intraventricular haemorrhage.

One hypothesis to explain the fluctuations in the velocity of cerebral blood flow is that swings in intrathoracic pressure, resulting from the combination of artificial and spontaneous respiration, change cardiac stroke volume and venous return to the heart. These then change arterial and venous pressures, which in turn affect cerebral blood flow.

This study aimed to examine the variability in cerebral blood flow velocity over the first 72 hours of life, particularly in relation to the pattern of interaction between spontaneous respiration and artificial ventilation.

Patients and methods

Infants, weighing less than 2500 g at birth who were admitted on the first day of life to the neonatal intensive care unit and receiving artificial ventilation were considered for the study. Infants who had an intraventricular haemorrhage detected by ultrasound scan at the time of the first study, or who had
a major congenital malformation were excluded, as was any infant admitted when one of the investigators was not available. Each infant was studied up to four times: at less than 12 hours of age and within six hours of 24, 48, and 72 hours of age. No further studies were made if the infant was paralysed, taken off the ventilator, or had sustained an intraventricular haemorrhage. During each study simultaneous recordings of blood flow velocity and respiratory interaction patterns were made by independent observers.

The variability of cerebral blood flow velocity was measured using a Duplex ultrasound scanner with range gated Doppler sampling (Advanced Technology Laboratories Mk 600). The Doppler sample volume was positioned near the anterior or middle cerebral artery using the real time image, and fine adjustments were made to give the loudest possible signal. To reduce bias the timing of recordings of blood flow velocity was chosen by the observer of the respiratory patterns who was unaware of the Doppler tracings. Each recording of blood flow velocity lasted seven seconds and comprised a paper printout of the fast Fourier transform of the Doppler frequency shift. The recordings were coded and analysed at a later date. The area within the peak frequency envelope for the whole (systole and diastole) of each of 10 cardiac cycles was measured by computer assisted planimetry (graphics tablet connected to a microcomputer). This measurement was made twice and the mean of the two measurements used. The mean error of area measurement (difference/mean×100) was 5.2%. The variability of blood flow velocity was calculated as the coefficient of variation of spectrum area for the 10 cardiac cycles in each record (CV=standard deviation/mean, expressed as a percentage).

The baby’s spontaneous respiration was monitored using a pneumatic respiration monitor (Graseby Dynamics MR10) taped just below the xiphisternum, as previously described. The ventilator circuit pressure was measured using a transducer (Mercury Electronics) connected near the top of the endotracheal tube. Signals were recorded on a polygraph (Gould 2600 series) at 25 mm/second.

The patterns of spontaneous respiration observed during ventilation were classified as apnoea, synchrony, or asynchrony. Apnoea indicated that the baby showed no spontaneous respiratory activity during the recording, synchrony that the baby’s inspiratory efforts were consistently in phase with the ventilator inflations, and asynchrony that the baby was breathing but with no regular association with the ventilator timing.

Six recordings were made on each occasion in one of two sequences: synchrony-asynchrony-synchrony or asynchrony-synchrony-asynchrony, two recordings being made during each phase. The sequence was chosen according to the pattern of interaction between the baby’s respiration and the ventilator cycles observed at the start of each period. The sequences were used to eliminate any time or order effects. If the baby was apnoeic only two recordings were made. Each phase lasted 5 to 10 minutes and the whole study period was less than 30 minutes.

Babies were induced to breathe asynchronously by using a ventilator rate of 30/minute with an inflation time of 0.5 seconds and deflation time of 1.5 seconds. They were induced to breathe synchronously by using ventilator settings chosen to match their spontaneous inspiratory and expiratory timing. The initial time settings for this were: rate 86/minute, inflation time 0.3 seconds, and deflation time 0.4 seconds. This was the average respiratory timing of a large comparable group of premature babies receiving ventilation who continued to breathe spontaneously. If necessary the time settings were altered to produce the best possible synchrony between baby and ventilator. At the end of each study, the infants were left breathing synchronously when possible. Details of blood gas measurements and ventilator settings were recorded.

Associations between variables were examined using Spearman’s test of non-parametric correlation. Comparisons between groups were made using the Mann-Whitney U test. The study was approved by the hospital ethical committee.

Results

Twenty infants with a mean birth weight of 1336 g and a mean gestational age of 29-2 weeks were studied on 42 separate occasions. Between one and four studies were made on each baby, less than four studies being made in some babies because of neonatal death (n=2), intraventricular haemorrhage (n=1), or stopping of mechanical ventilation (n=15). The table shows the general information about the babies at the time of the studies.

<table>
<thead>
<tr>
<th>Measurement</th>
<th>Median</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gestation (weeks)</td>
<td>29</td>
<td>26-39</td>
</tr>
<tr>
<td>Birth weight (g)</td>
<td>1336</td>
<td>630-1876</td>
</tr>
<tr>
<td>Inspired oxygen (%)</td>
<td>50</td>
<td>25-90</td>
</tr>
<tr>
<td>Peak pressure (kPa)</td>
<td>1.96</td>
<td>1.27-2.94</td>
</tr>
<tr>
<td>pO2 (kPa)</td>
<td>8.5</td>
<td>5-19</td>
</tr>
<tr>
<td>pCO2 (kPa)</td>
<td>5.0</td>
<td>3-8</td>
</tr>
<tr>
<td>pH</td>
<td>7.36</td>
<td>7.2-7.5</td>
</tr>
</tbody>
</table>
Fig 1  Median variability in cerebral blood flow velocity and interquartile ranges for each study age. *p<0.0001 [v. less than 12 hours].

Fig 2  Example of blood flow velocity and respiratory recordings for a baby during asynchronous breathing. There is no regular correspondence between spontaneous respiratory pattern and ventilator. Note variability in waveform area.

Fig 3  Example of blood flow velocity and respiratory recordings for a baby during synchronous breathing. Note reduced variability in waveform area.

No association was found between blood flow velocity variability and birth weight, gestational age, peak ventilator pressure, arterial oxygen or carbon dioxide tensions. There was a weak but significant association with pH, increasing pH correlating with increasing variability (r=0.23, p=0.001).

Of the 20 infants in the study three developed an intra ventricular haemorrhage. The predominant pattern of interaction at the onset of each study period was split equally between synchrony and asynchrony in these patients. There were no characteristic features to the waveform pattern of cerebral blood flow velocity in these babies.
The variability in cerebral blood flow velocity of premature babies receiving assisted ventilation was fairly high within 12 hours of birth and fell progressively over the first 48 hours of life. This change was seen during recordings made during both synchronous and asynchronous ventilation (Fig 5). These results suggest that variability is greatest at the time when intraventricular haemorrhage usually occurs. Drayton and Skidmore suggested that the major intracranial arteries are probably dilated shortly after birth, and that they constrict over the first 48 hours of life in response to rising blood pressure. These authors hypothesised that while the cerebral arteries are dilated they are less efficient at damping arterial pressure surges and thus place the premature infant at particular risk of intraventricular haemorrhage in the early hours of life.

A second explanation for our results is that systemic blood pressure may have fluctuated less after 48 hours of life and that this (rather than any change in the diameter of the cerebral vessels) was responsible. Unfortunately systemic variability of arterial blood pressure could not be monitored as the lumens of the umbilical arterial catheters were too narrow and had side rather than end openings. These catheters result in severe damping of any pressure fluctuations.

This study shows that the pattern of interaction between spontaneous respiration and mechanical ventilation influences the variability of the velocity of blood flow in the major intracranial arteries: apnoea and synchronous breathing result in low variability and asynchronous breathing in significantly higher variability. It is not possible to comment on the association between the pattern of ventilation interaction and the occurrence of intraventricular haemorrhage in our small study.

Theoretically, changes in the velocity of blood flow could result either from changes in blood flow or in the calibre of arteries, but the calibre of a vessel is unlikely to change rapidly enough to produce variations in velocity from beat to beat. Experimental evidence exists that suggests that the cerebral vessels with a diameter of over 200 μm that change in calibre during autoregulation at physiological blood pressures take between three and seven seconds to change size. It seems most likely that the variations from beat to beat in velocity of the blood flow seen in this study represent actual changes in blood flow. The effects of different patterns of interaction between spontaneous respiration and mechanical ventilation on venous return and cardiac output provide a logical explanation for this. In a subject breathing normally, respiration affects the circulation. During inspiration the intrathoracic pressure falls and blood is drawn into the right side of the heart from the systemic venous circulation leading to a rise in right heart output. At the same time, because the fall in intrathoracic pressure causes dilatation of the pulmonary venous system, return of blood to the left side of the heart is reduced and left heart output falls; this causes a drop in systemic blood pressure. The fluctuations in venous return and arterial blood pressure are fairly small in the healthy subject, but they may be exaggerated by the airways obstruction and hypovolaemia that may occur in sick babies receiving assisted ventilation.
When a baby is subject to asynchronous ventilation the changes in intrathoracic pressure are likely to be greater and more irregular. During synchrony the intrathoracic pressures and flows in systemic venous and arterial blood pressure will occur when the baby breathes in during the deflation phase of the ventilator, and high intrathoracic pressures (and rises in systemic venous and arterial pressure) will occur when the baby breathes out while the ventilator is attempting to inflate the lungs. This rise is particularly likely if the baby actively expires. If a baby breathes synchronously with the ventilator, then spontaneous breathing in coincides with inflation and the change in intrathoracic pressure is small. Asynchronous breathing would thus result in more erratic and larger fluctuations in arterial blood pressure and in venous drainage from the head than synchronous breathing or apnoea.

An increased variation in occipitofrontal circumference during positive pressure ventilation has been reported that presumably reflects changes in cerebral blood volume. Variations in cerebral venous and arterial velocities measured with Doppler ultrasonography also occur in phase with artificial ventilation, and a reduction in peak inspiratory pressure or discontinuation of ventilation reduced the variability in velocity. The greater variability in cerebral blood flow velocity in the early hours of postnatal life could partly explain the usual timing of periventricular haemorrhage.

These results show that asynchronous ventilation leads to greater fluctuations in the velocity of blood flow through the intracranial arteries. If, as suggested by other authors, these fluctuations increase the risk of intraventricular haemorrhage, then measures to reduce the amount of asynchronous ventilation might reduce this risk.

We thank Professor JA Davis, Dr NRC Robertson, and Dr G Gandy for advice and encouragement, and the junior medical and nursing staff for their tolerance and care of the patients. This study was supported by the University of Cambridge Baby Research Fund, Children Nationwide, the Trustees of Addenbrooke’s Hospital, and the East Anglian Regional Health Authority.

References

7 Greenough A, Wood S, Morley, Davis JA. Pancuronium prevents pneumothoraces in ventilated preterm babies who actively expire against positive pressure inflation. Lancet 1984;i:1-3.

Correspondence and requests for reprints to Dr JM Rennie, Department of Paediatrics, University of Cambridge, Level E8, Addenbrooke’s Hospital, Cambridge CB2 0QQ.

Received 20 July 1987