Hypocapnia and cerebral ischaemia in hypotensive newborn piglets

A Whitelaw, B R Karlsson, K Haaland, I Dahlin, P A Steen, M Thoresen

Abstract
This study tested the hypothesis that hypocapnia superimposed upon hypotension produces a further reduction in cerebral blood flow velocity (CBFV). In 12 newborn piglets, CBFV was measured continuously by Doppler ultrasound through an artificial fontanelle. Hypotension was induced by removing 30 ml/kg of blood. Increasing the ventilator rate reduced the average arterial carbon dioxide tension from 5-5 to 2-0 kPa. When mean arterial pressure (MAP) was held steady at 45 mm Hg or above, hypocapnia produced a substantial drop in CBFV, but in all the piglets with MAP below 38 mm Hg, hypocapnia failed to change CBFV by 10%. Hypocapnia produced an increase in lactate in sagittal sinus blood but cerebral venous hypoxanthine concentrations were not affected by hypocapnia. Hyperventilation (without haemorrhage) produced a significant drop in MAP, preventable by infusing colloid.

Hypocapnia itself does not further reduce CBFV in the hypotensive piglet. However, the pressure effect of hyperventilation may significantly impair the cerebral circulation.

Despite the great improvements in the survival of premature infants, there are reports of increasing numbers of handicapped survivors with cerebral palsy.1 2 The most important neuropathological lesion associated with spastic diplegia in premature infants is periventricular leucomalacia. This is a bilateral cerebral lesion in the white matter furthest from the arterial blood supply. The aetiology is not fully understood and no treatment has been shown to be effective in preventing periventricular leucomalacia. On theoretical grounds, it is likely that hypotension is an important causative factor but the occurrence of periventricular leucomalacia is not consistently associated with hypotension.3 There must be other aetiological factors.

It has been proposed that a very low arterial carbon dioxide tension (Paco2) (hypocapnia) may be important in the aetiology of ischaemic brain damage in newborns.4 When blood pressure is normal, raising Paco2 increases cerebral blood flow and decreasing Paco2 decreases cerebral blood flow. When blood pressure falls, the brain tries to maintain blood flow by progressively vasodilating and eventually becomes maximally vasodilated. If hypocapnia caused cerebral vasoconstriction in this situation, cerebral blood flow could be reduced to a critically low level with ischaemic damage following. Although brain lesions have not been demonstrated in experimental animals after hypocapnia,5 slowing of the electroencephalogram has been noted in adult humans during hyperventilation. These changes could be reversed by hyperventilating with 100% oxygen.6 The normal escape mechanisms that inhibit excessive vasoconstriction may not operate in the immature neonatal cerebral circulation.

Inadvertent hypocapnia may occur when premature infants are mechanically ventilated for transport by ambulance or helicopter from the place of birth to the neonatal intensive care unit as infants do not usually have any form of carbon dioxide monitoring during transport. Hypocapnia may occur at any time that an infant is mechanically ventilated without careful monitoring of Paco2, for example during surgical anaesthesia. This is particularly likely with infants who do not have respiratory distress syndrome and stiff lungs.

Calvert et al reported a case-control study of 15 infants with periventricular leucomalacia.7 The infants with periventricular leucomalacia had experienced significantly lower Paco2 values during the first 72 hours than had the control infants. The length of time that Paco2 was below 3-3 kPa was longer in the infants who subsequently developed periventricular leucomalacia. Calvert et al had no data on blood pressure but discussed the likelihood that the infants with periventricular leucomalacia had suffered hypotension as well as hypocapnia. Evidence for the involvement of hypocapnia and hypoperfusion in the aetiology of neurological deficit after cardiopulmonary bypass in adult patients has also been reported.8

In this study we tested the hypothesis that hypocapnia superimposed upon hypotension in the newborn can further reduce cerebral blood flow velocity and thereby increase the risk of cerebral ischaemia. None of the previous papers on hypocapnia, hypotension, and cerebral blood flow have dealt with newborns and the study would be unethical in human infants. The newborn piglet was chosen as the experimental model because of the great similarities in cardiovascular and respiratory systems, cerebral circulation, and body size between the piglet and human infant.

Methods
The experiments were performed on 12 newborn piglets aged from 12 hours to 3-5 days and weighing 1-1 to 2-8 kg, which were kept with the sow until three hours before the

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Hypocapnia

With the rate of HYPOTENSION hyperventilation while was been through back expiratory animals Three with ventilator from fontanelle was measured by standard laboratory recording mean Median and sutures meet.

A tracheostomy was performed and each piglet was paralysed with pancuronium 0-2 mg/kg intravenously and mechanically ventilated with 25–30% oxygen during surgery, care being taken to avoid any hypoxaemia. Both umbilical arteries and the umbilical vein were catheterised in piglets less than 48 hours old. In slightly older piglets a femoral artery and subclavian vein were catheterised. These catheters were used for blood sampling, continuous arterial pressure recording, and infusion of fluids and drugs. In five animals a catheter was inserted into the right internal jugular vein with the tip just below the brain and in two animals a catheter was inserted into the sagittal sinus. These catheters were used for hypoxanthine and lactate blood sampling. Blood samples were separated within one minute and plasma kept frozen until hypoxanthine was measured by high precision liquid chromatography. Plasma lactate was measured by standard laboratory techniques. In order to measure intracerebral blood velocities with Doppler ultrasound, an artificial fontanelle of 1 cm diameter was made just lateral to the midline where the cranial sutures meet.

All values are given as median values with the 95% confidence interval and are listed in the table. Comparisons of grouped data were made using the Wilcoxon signed rank test.

The experimental protocol was as follows:

HYPOCAPNIA DURING NORMOTENSION

Median mean arterial pressure (MAP) at the start of the protocol was 48 mm Hg (37 to 75). After recording cerebral blood flow velocity (CBFV) and MAP continuously for five minutes with a normal PaCO₂ (5-5 kPa), hyperventilation was produced by increasing the frequency of the ventilator from 20 (inspiratory time 0-3 seconds, expiratory time 2-7 seconds) to 100 breaths/minute (bpm) (inspiratory time 0-3 seconds, expiratory time 0-3 seconds). This produced a rapid fall in PaCO₂ within 10 minutes to 2-0 kPa. Three animals received 20 ml/kg 5% albumin during hyperventilation to correct the tendency for MAP to fall. CBFV recording was continued while hyperventilation was continued for a further 10 minutes and then normal ventilation was resumed (18 to 20 bpm). The PaCO₂ rose back to the normal range.

HYPOCAPNIA DURING HYPOPTENSION

With the PaCO₂ at 5-4 kPa, blood was withdrawn through a catheter into heparinised syringes at a rate of 1–2 ml/kg/minute until 30 ml/kg had been removed. This produced a median fall in MAP of 37% to 32 mm Hg as well as a median fall of 35% in CBFV.

HYPOTENSIVE HYPOCAPNIA

In a stable hypotensive condition, at least 30 minutes after the end of the bleed, baseline recordings with a normal PaCO₂ were obtained. Hyperventilation was again produced by ventilating at 100 bpm. The PaCO₂ fell from 5-5 kPa to 1-9 kPa. Blood was transfused back into the piglet during hyperventilation at a rate sufficient to hold the MAP steady. Hyperventilation and recording of CBFV and MAP continued. After 20 minutes of hyperventilation the last blood samples were obtained and the experiment was terminated by rapid injection of potassium chloride.

CBF RECORDINGS

As long as the cross sectional area of the vessel recorded from is constant, changes in blood velocity are proportional to changes in blood flow. A 10 MHz Doppler ultrasound system (SD 100 Vingmed Sound) was used. By adjusting the depth of the sample volume and direction of insonication through the artificial fontanelle, precise localisation on an intracerebral artery could be achieved. In each piglet, the same artery (1-7 cm deep and 1-5 cm lateral to the midline) was used for recording at a constant angle with the transducer held in place manually. The Doppler system was interfaced to an Apricot XI computer. Every 5 ms the computer was fed the mean CBFV calculated from the Doppler spectrum, MAP, and heart rate. CBFV, MAP, and heart rate were all displayed in real time on a screen during the whole experimental period and stored on disc.

Results

HYPOCAPNIA DURING HYPOPTENSION

Hyperventilating the piglets by increasing the ventilator rate from 20 to 100 bpm produced an appreciable drop in MAP. This is shown in the table and illustrated in fig 1. MAP in nine piglets fell from a median of 48 mm Hg (37–75) to a median of 36 mm Hg (29–56) and in three piglets where albumin was given, MAP fell from a median of 46 to 41 mm Hg. Thus during hyperventilation the median MAP of all 12 piglets was 41 mm Hg (33 to 51). In all the piglets with a MAP above 44 mm Hg there was a definite fall in cerebral blood flow velocity during hyperventilation. These young piglets nearly all showed a proportional decrease in CBFV with MAP. Hence when calculating the effect of low PaCO₂ on CBFV, the effect of blood pressure change has to be corrected for. This is illustrated in fig 1 where CBFV fell by nearly 50% whereas MAP fell by less than 25% during hyperventilation. Thus 25% was subtracted from the initial CBFV to calculate the starting value from which the change during hypocapnia was calculated. The table shows the CBFV fell by a median of 20% (10–33) during this normotensive hypocapnia. A cerebrovascular response to hypocapnia was considered to be present if CBFV fell by at least 10%.
Cardiovascular changes during experiment. Values are medians (95% confidence intervals)

<table>
<thead>
<tr>
<th></th>
<th>Normotensive hypocapnia</th>
<th>Haemorrhagic hypotension</th>
<th>Hypotensive hypocapnia</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Before</td>
<td>During</td>
<td>Before</td>
</tr>
<tr>
<td>MAP (mmHg):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(n=9)</td>
<td>48(47 to 75)</td>
<td>36(29 to 56)</td>
<td>51(46 to 56)</td>
</tr>
<tr>
<td>(n=3, albumin)</td>
<td>46(45 to 55)</td>
<td>41(40 to 51)</td>
<td>190(165 to 220)</td>
</tr>
<tr>
<td>(n=12)</td>
<td>48(46 to 68)</td>
<td>41(33 to 51)</td>
<td>134(101 to 210)</td>
</tr>
<tr>
<td>Heart rate (bpm)*</td>
<td>178(167 to 198)</td>
<td>195(180 to 210)</td>
<td>178(167 to 198)</td>
</tr>
<tr>
<td>PaO2 (kPa)*</td>
<td>14.6(12.8 to 17.0)</td>
<td>13.9(11.7 to 16.0)</td>
<td>14.0(12.4 to 16.7)</td>
</tr>
<tr>
<td>PacO2 (kPa)*</td>
<td>5.5(5.0 to 6.0)</td>
<td>2.0(1.7 to 2.4)</td>
<td>5.4(5.1 to 6.0)</td>
</tr>
<tr>
<td>pH*</td>
<td>7.36(7.32 to 7.40)</td>
<td>7.70(7.65 to 7.76)</td>
<td>7.35(7.35 to 7.41)</td>
</tr>
<tr>
<td>Base excess (mmol/l)*</td>
<td>-2.7(-4.1 to -0.9)</td>
<td>1.0(-0.9 to 5.1)</td>
<td>-2.1(-5.1 to 0.0)</td>
</tr>
<tr>
<td>% Reduction in CBFV*</td>
<td>20(10.0 to 33.0)</td>
<td>34.5(17.0 to 41.0)</td>
<td>0.0(0.0 to 6.0)</td>
</tr>
</tbody>
</table>

* n=12.

Figure 1 Continuous recording of CBFV and MAP in a piglet during hyperventilation with normal blood pressure. There is a fall in MAP but a much larger fall in CBFV indicating a cerebrovascular response to hypocapnia.

Figure 2 Continuous recording of CBFV and MAP in a piglet during hyperventilation with hypotension after haemorrhage. Blood was transfused back to maintain a steady blood pressure during hyperventilation. There was no change in CBFV indicating an absent cerebrovascular response to hypocapnia.

HYPOCAPNIA DURING HAEOMORRHAGIC HYPOTENSION

During hyperventilation after the haemorrhage, MAP was maintained by blood retransfusion. In eight out of nine piglets with MAP below 38 mm Hg, hypocapnia failed to produce any further fall in CBFV (fig 2), and the ninth piglet showed only a 6% fall. In one piglet with MAP

CEREBROVASCULAR RESPONSE TO HYPOCAPNIA AT DIFFERENT MAP

When all 24 hyperventilation challenges are considered together as in fig 4, it can be seen that the cerebrovascular response to hypocapnia was clearly present when the MAP was over 44 mm Hg. When MAP was below 38 mm Hg, CBFV responses were absent or less than 10%.

Figure 5 shows that jugular venous or sagittal sinus plasma hypoxanthine concentration showed no significant increase during either normotensive hypocapnia or hypotensive hypocapnia. In the seven animals examined there was an increase in plasma hypoxanthine as the
Hypocapnia or during hemorrhagic hypotension

Figure S
Serial plasma hypoxanthine concentrations in blood draining the brain, either from the jugular veins (five piglets) or from the sagittal sinus (two piglets). Numbers on X-axis indicate samples before (1) and during (2) normotensive hypocapnia, before (3) and during (4) hemorrhagic hypotension and before (5) and during (6) hypotensive hypocapnia. There was no change in plasma hypoxanthine during either normotensive (2 vs 1) or hypotensive (5 vs 5) hypocapnia. There was a significant increase in hypoxanthine after the hemorrhage (5 vs 3, p<0.005).

Discussion
When MAP was steady at a level below 38 mm Hg, hypocapnia produced negligible reduction in CBFV. This is clear evidence against our hypothesis that hypocapnia could further impair the cerebral circulation during hypotension. It would be an appropriate physiological response for the cerebral circulation to vasodilate in response to hypotension and to try to maintain the best possible cerebral circulation at all costs.

Thus it seems that the cerebral vasodilator response to hypotension cannot be over-ridden by the cerebral vasoconstrictor response to hypocapnia. Hypoxanthine is probably the best available plasma marker of cellular hypoxia (or failure to make ATP) and the lack of any significant increase in hypoxanthine in jugular or sagittal sinus blood is further evidence that hypocapnia does not, by itself, produce cerebral ischemia.

The increase in lactate in sagittal sinus blood is not necessarily evidence of cerebral ischemia. There is much evidence that the increase in pH with hyperventilation changes the activity of glycolytic pathway enzymes and thus increases lactate production while cerebral levels of phosphocreatine, ATP, and energy charge remain unchanged.14 Mitchenfelder and Theye have demonstrated in dogs that hypocapnia can shift the oxygen-haemoglobin dissociation curve in sagittal sinus blood to the left (Bohr effect) thus reducing oxygen availability to the tissues.15 This was associated with a change towards anaerobic metabolism. Lactate has been postulated to have a role as a cerebral vasodilator but the rise in lactate during hypotensive hypocapnia was not accompanied by further vasodilatation (fig 2). This finding suggests that the cerebral circulation was already maximally vasodilated during hypotension.

Do our findings mean that overventilation is harmless? Not necessarily. The anecdotal observations of an association between hypocapnia and subsequent neurological impairment may be due, not to a direct effect of hypocapnia on the cerebral circulation, but to the pressure effect of hyperventilation on the circulation. We observed that hyperventilation produced a median drop in MAP of 12 mm Hg. A fall in blood pressure of such magnitude combined with the Bohr effect on oxygen-haemoglobin dissociation could certainly impair cerebral oxygenation in a critically ill infant. We have previously described how increases in ventilator pressure can reduce CBFV in newborn infants.16

If the infant's lungs are relatively compliant the ventilator pressure will be transmitted to the circulation thus producing a resistance to venous return to the right atrium, which would tend to reduce cardiac output. In addition, increased jugular venous pressure constitutes a resistance to cerebral blood flow.

In conclusion, the cerebral circulation of the hypotensive newborn piglet does not vasocostrict in response to hypocapnia but positive pressure hyperventilation may significantly impair the cerebral circulation and oxygenation. Thus it would seem wise to avoid hypocapnia in mechanically ventilated infants particularly those with hypotension, hypovolaemia, or anaemia.

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