Intravenous aminophylline and cerebral blood flow in preterm infants

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Abstract
The effect of aminophylline on cerebral blood flow (CBF) was studied in 10 preterm infants who were receiving 6-2 mg/kg intravenously over 20 minutes followed by a maintenance infusion. CBF was measured intermittently using near infrared spectroscopy. Heart rate, blood pressure, oxygen saturation, and transcutaneously measured carbon dioxide tension (TcPCO$_2$) were recorded continuously. Aminophylline administration was associated with a fall in CBF from a median of 15-9 ml/100 g/min to 11-2 ml/100 g/min. Median fall in CBF was 4-1 ml/100 g/min (95% confidence interval 1-7 to 6-5). Heart rate rose and TcPCO$_2$ fell in all infants, median fall being 0-66 kPa. The reduction in CBF was greater than would be expected on the basis of the modest fall in TcPCO$_2$.

Aminophylline and other methylxanthines are frequently used in the neonatal period and have been reported to cause reduction in cerebral blood flow.$^1$ We have used near infrared spectroscopy (NIRS) to study cerebral haemodynamics and oxygenation before and after intravenous loading of preterm infants with aminophylline, according to the usual regimen for treatment of apnoea of prematurity on our unit.

NIRS is a non-invasive optical technique which provides continuous outside measurement of the amounts of oxyhaemoglobin, reduced haemoglobin, and oxidised cytochrome aa3 in the light path between two optodes applied to either side of the infant's head.$^2$ Each of these three chromophores has a characteristic absorption spectrum for near infrared light. Light at six discrete wavelengths is transmitted from laser diodes to the head through fiberoptic cables. A sensitive photomultiplier detects the small amounts of light transmitted across the head, and the attenuation at different wavelengths can be used to calculate chromophore concentrations.

A transient increase in the infant's inspired oxygen concentration will cause a bolus of oxyhaemoglobin to enter the brain at a rate which is determined by cerebral blood flow (CBF) and can be measured by NIRS. CBF can be quantified in this way, using the method described by Edwards et al.$^3$

Patients and methods
Ten preterm infants (gestation 23–31 weeks, median 26) were studied at 4–28 (median 9) days of age. Median birth weight was 830 g, range 519–2056 g. Seven of the infants were receiving mechanical ventilation, one was in headbox oxygen, and two were being treated with continuous positive airways pressure.

Aminophylline was prescribed for the treatment of apnoea of prematurity or to assist weaning from ventilatory assistance. A loading dose of 6-2 mg/kg body weight was given intravenously over 20 minutes and was followed by a maintenance infusion of 4-4 mg/kg/day. This regimen was recommended by Jones and Bailie on the basis of pharmacokinetic studies, and was calculated to give a serum concentration of 28–66 µmol/l one hour after loading.$^4$

Oxygen saturation was measured by a pulse oximeter on the right hand working in beat to beat mode, and mean arterial blood pressure was measured through an umbilical or peripheral arterial cannula using a Hewlett-Packard pressure transducer. Heart rate, transcutaneous carbon dioxide tension (TcPCO$_2$), oxygen saturation, and mean arterial blood pressure were monitored continuously throughout the 2–4 (median 3) hour period of each study. Physiological measurements were stored on disc at 2–20 second intervals using the data storage system of the NIRS spectrometer.

The near infrared measurements were made using the NIR-1000 (Hamamatsu Photonics KK) spectrometer. Optodes were applied to each temporal region, kept in place by an elasticated bandage, and covered by a light occluding hat which prevented contamination of the near infrared signal by external light. CBF was measured using a change in oxyhaemoglobin as a tracer. By the Fick principle, CBF can be calculated as the ratio of oxyhaemoglobin accumulated in the brain to the quantity introduced into the brain during a time period less than cerebral transit time (which is approximately 10 seconds). After a period when the oxyhaemoglobin signal and oxygen saturation were stable (with a saturation of 88–96%), the inspired oxygen concentration was suddenly increased in order to induce a transient increase in saturation of 5–8%. Cerebral oxyhaemoglobin and oxygen saturation were recorded at two second intervals during this manoeuvre for subsequent calculation of CBF.

CBF measurements were made on one to four (median three) occasions during the 135 minutes before drug infusion, and on one to five (median three) occasions in the 110 minutes after drug infusion. All infants were clinically stable during the period of the study. CBF values were not available 'on line' during the recording period. The near infrared and physiological data...
stored on disc in the spectrometer were transferred to a microcomputer spreadsheet for quantitative analysis at a later time. The data recorded during each brief period of induced rise in oxygen saturation were inspected, and all episodes in which data fitted a prospectively determined set of criteria were analysed and a CBF measurement determined. Measurements of physiological variables were taken during the periods of CBF measurement and later used in statistical analysis. Mean values of readings from each infant in the period before the drug were compared with mean values from the period after the drug was given using Wilcoxon signed ranks test for paired non-parametric data.

Results

Median CBF in the predrug period was 15.9 ml/100 g/min (95% confidence interval (CI) 13.5 to 17.7) and median CBF in the postdrug period was 11.2 ml/100 g/min. Eight infants showed a fall in CBF, one infant no change and one showed a 0.9 ml/100 g/min (6.9%) rise. The median fall in CBF was 4.1 ml/100 g/min (95% CI 1.7 to 6.5), or 28.9% (95% CI 12.3 to 37.7). Figure 1A shows the individual data points. The fall in CBF was significant on paired testing at p<0.01.

Figure 1B shows that heart rate rose in all infants (p<0.01), by a median of 12 beats/min (95% CI 9 to 19.2).

Figure 1C shows that a fall in TcPCO2 was demonstrated in all infants (p<0.01). Median fall was 0.66 kPa (95% CI 0.40 to 0.84). There was no consistent change in mean arterial blood pressure after aminophylline infusion (fig 1D).

If the changes in CBF (dCBF) after aminophylline were completely due to the changes in carbon dioxide tension (dPCO2), then a quantitative relationship between dCBF and dPCO2 would be expected. However, there was no significant correlation between dCBF and dPCO2, either expressed in absolute terms (correlation coefficient r=0.57, n=10, NS) or as a fraction of resting values (r=0.49, n=10, NS).
**Discussion**

Aminophylline and other methylxanthines are commonly used for the treatment of apnoea of prematurity, and to assist weaning of low birthweight infants from mechanical ventilation. Preterm infants treated with methylxanthines show increased alveolar ventilation, decrease in the incidence of apneic episodes, and higher cardiac output due to increases in both heart rate and stroke volume. These diverse effects are attributed to adenosine receptor antagonism, as very little phosphodiesterase inhibition occurs at therapeutic serum concentrations. As adenosine is known to be a cerebral vasodilator, aminophylline may be expected to cause vasoconstriction and reduction in CBF.7

Wechsler et al in 1950 were the first group to show reduction in CBF after aminophylline administration.8 Using the Kety-Schmidt method they showed a 25% fall in CBF of adults receiving an intravenous loading dose. The effect of methylxanthines on cerebral blood flow velocity (CBFV) of preterm infants has been measured by Doppler ultrasound in four studies. Rosenkranz and Oh showed a 17% reduction in CBFV in two hours after a 5 mg/kg bolus of aminophylline9 which they attributed to a concomitant fall in Pco2. Ghai et al found no change in Pco2 or CBFV in 13 preterm infants studied 50 hours after starting enteral therapy.10 Two recent studies have shown no effect of systemic doses of caffeine on CBFV in preterm infants,11,12 although one of these studies11 showed a significant fall in Pco2. However, Doppler studies are difficult to interpret because CBFV may not reliably reflect CBF in this clinical situation when heart rate, blood pressure, Pco2, and cerebral vessel diameter may all be changing. Pr dys et al, using intravenous xenon13 clearance to measure CBF, reported a 13%-8% reduction in CBF but unaltered visual evoked responses, one hour after 10 mg/kg intravenous aminophylline.1

NIRS can be used to determine cerebral blood flow with minimal patient disturbance. Since the description of the technique1 there have been two reports validating the results of the NIRS technique against xenon clearance methods.13,14 Both NIRS and xenon clearance give surprisingly low values for CBF in the newborn, but they are consistent with values reported by Volpe et al using positron emission tomography.15

Nine infants in this present study had two or more measurements of CBF before the dose of aminophylline and the mean coefficient of variation of CBF measurement during this clinically stable period was 19% (95% CI 12.8-25.9). This coefficient is high and presumably reflects, at least in part, methodological limitations, but does not alter the significance of the changes in CBF seen on paired testing after aminophylline.

There was a small but consistent fall in Pco2, but we do not consider this to be the full explanation for the considerable fall in CBF. Reported values for carbon dioxide reactivity of the cerebral circulation of preterm infants vary considerably. Pr dys et al recently showed, using intravenous xenon13, a 28-9% change per kPa change in Pco2.16 Using this value the fall in Pco2 we have reported in this study would be expected to result in a fall in CBF of 19%, whereas a median decrease of 29% was actually observed. In addition, if decreased Pco2 was the sole mechanism for the fall in CBF, there would be some relationship between the extent to which Pco2 and CBF fall in individual infants, which was not demonstrated. It appears, therefore, that an intravenous loading dose of aminophylline leads to a fall in CBF which cannot be entirely explained by a small decrease in Pco2.

A decrease of CBF, and hence cerebral oxygen delivery, will generally be compensated for by increased oxygen extraction resulting in a lower cerebral venous oxygen tension. There may be no clinical sequela to considerable falls in CBF such as shown here with aminophylline treatment, or previously reported with indomethacin.13 However, if cerebral oxygen delivery is already compromised in a sick preterm infant with hypotension and impaired oxygenation, further reduction of CBF of the magnitude reported in this study could potentially have serious sequela.

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