Cerebral distribution of cardiac output in newborn infants

T Kusaka, K Okubo, K Nagano, K Isobe, S Itoh

Cerebral blood flow and output of the left ventricle were simultaneously investigated in 17 infants using multichannel near infrared spectroscopy and pulse dye densitometry with indocyanine green. Cardiac output and cerebral blood flow were positively related. The control of cardiac output is important in the regulation of cerebral blood flow in infants.

The major regulatory mechanisms for cerebral blood flow (CBF) in infants are autoregulation, arteriolar CO2, O2 delivery, blood glucose, and neural activity. It has also been shown that the cardiovascular system regulates CBF through variation in cardiac output (CO) and distribution of blood flow.

A new method of calculating CBF using indocyanine green (ICG) as a tracer with near infrared spectroscopy (NIRS) has been developed.1,2 We previously reported that multichannel blood flow through variation in cardiac output (CO) and distribution of CBF can be used to measure regional CBF distribution in infants.2  NIRS (MNIRS) and pulse dye densitometry (PDD) using ICG can be used to measure regional CBF distribution in infants.2  PDD uses the general principles of pulse oximetry to measure the concentration ratios of ICG and haemoglobin (Hb) in arterial blood. Therefore ICG concentration in arterial blood can be estimated without collecting blood if Hb concentrations are obtained in a clinical setting.1,3

In this study, measurement of CBF using MNIRS and PDD and measurement of CO using PDD with ICG were carried out simultaneously in 17 infants, and the relation between CO and CBF was investigated.

SUBJECTS AND METHODS

Measurements were carried out in 19 infants undergoing neonatal intensive care at the Maternal Perinatal Center of Kagawa University Hospital. Written informed consent was obtained from the parents of each infant. The study was also approved by a local ethics committee. Data from two infants were excluded from the analysis because the measurements were affected by movement artefacts. Successful measurements were obtained in 17 infants (mean (SD) gestational age 32.9 (4.3) weeks (range 24.1–38.1) and birth weight 1588 (686) g (range 651–3200)). Median age at the time of measurement was 5 days (range 0–82).

A bolus of ICG (0.2 mg/kg) was injected into the peripheral vein of each infant. Eight light incident and eight detector fibres were placed on the right or left parietotemporal region of the head, each with an interoptode distance of 2 cm, and 24 measuring positions were defined. Changes in cerebral ICG concentration in a 6 cm × 6 cm field were recorded using MNIRS (Optical Encephalography System; Hitachi Medical Co Ltd, Chiba, Japan). Data were collected every 0.1 second after the appearance of dye in the optical field. The accumulation of cerebral ICG at a specific time later (Q(t)) was calculated. Simultaneously, PDD (DDG-2001; Nihon-Koden Co, Tokyo, Japan) was used to measure the arterial blood concentration of ICG (Pa). The optical probe was attached to the right or left wrist. CBF was calculated from the Fick equation modified for zero venous concentration during the first six seconds after the appearance of dye in each optical field. With the use of measured cerebral ICG (mg/l/cm), arterial ICG (mg/l), measured interoptode distance (cm), and differential path length factor for infants (4.39), where K is a constant reflecting the cerebral tissue density (1.05), and correcting to 100g tissue mass, CBF can be obtained as follows:

\[
\text{CBF (ml/100 g/min)} = \frac{K \times Q(t)}{D (mg)} \times \text{P}_{\text{a}}
\]

CBF was defined as the mean from 24 channels.

Left ventricle CO was calculated as follows from the first dilution curve on a PDD:

\[
\text{CO (l/min)} = \frac{D (mg)}{G (mg/min/l)}
\]

where D is the amount of ICG administered, and G is the area under the first dilution curve obtained by extrapolation of the first decay slope to infinity.

RESULTS

Table 1 shows the values for left ventricle CO and CBF. CO and CBF in the parietotemporal region were 202.8 (90.6) ml/kg/min and 15.3 (4.2) ml/100 g/min respectively. A significant positive relation was found between CO and CBF: CBF = 0.03CO + 8.71 (r = 0.70, p = 0.002) (fig 1). The relation between CBF and mean arterial blood pressure or between CBF and PvCO2 was not significant. However, CBF and blood Hb showed a significant positive relation (r = 0.49, p = 0.045).

DISCUSSION

This is the first report of a significant positive relation between CO and CBF in infants. The results of this study show that CO is a major factor affecting CBF. From our data,

![Figure 1](http://fn.bmj.com/)

**Figure 1** Correlation between cardiac output and cerebral blood flow measured by multichannel near infrared spectroscopy and pulse dye densitometry using indocyanine green. The linear regression line is described by CBF = 0.03CO + 8.71, with a correlation coefficient of 0.70 and p = 0.002.

**Abbreviations:** CBF, cerebral blood flow; CO, cardiac output; Hb, haemoglobin; ICG, indocyanine green; MNIRS, multichannel NIRS; NIRS, near infrared spectroscopy; PDD, pulse dye densitometry.

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the blood flow distribution to the brain of left ventricular output is estimated to be 8–11% if the brain is assumed to weigh 10–15% of body weight. This value is higher than the combined CO reported for the fetus of a lamb (3%). This is because cerebral O2 consumption and metabolic rate in infants are higher than in the fetus of a lamb.

Measurements of CO and CBF using MNIRS and PDD are simple and fast, and repeated measurements in an infant at the bedside are possible. There is no irradiation and no need for blood collection, and measurements can be made with little stress to the patient. However, further studies are needed in infants, especially for evaluation of the sensitivity and reliability of CO and CBF measurements using MNIRS and PDD, before the method can be applied to a clinical setting.

In conclusion, we found this procedure both feasible and very useful for the elucidation of CO and CBF disturbance during the acute phase of illness in infants.

Competing interests: none declared

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Accepted 2 August 2004

REFERENCES


Table 1

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MABP, Mean arterial blood pressure.
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Arch Dis Child Fetal Neonatal Ed 2005 90: F77-F78
doi: 10.1136/adc.2004.058487

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