Postnatal evolution of slow variability in cerebral blood flow velocity

Heather Coughtrey, Janet M Rennie, David H Evans

Abstract

The evolution of low frequency variability in cerebral blood flow velocity (CBFV) was examined in 30 consecutive admissions of very low birthweight babies. Measurement of CBFV was made on the first day of life and at weekly intervals until discharge. Altogether 133 recordings were subjected to visual classification and described as showing presence or absence of variability at a frequency between 1 and 5/min. Amplitude of variability was expressed as the largest variation in peak systolic velocity as a percentage of the maximum systolic change.

Slow variability was usually obvious and was sometimes present for only part of the minute recorded. There was a significant trend for decreased incidence of slow variability in relation to both postconceptional and postnatal ages. Amplitude of slow variability was also damped with increasing age.

Slow variability appeared to be unrelated to the type of sedation, severity of illness, or intracranial pathology; it may be a normal phenomenon in which damping occurs as the autonomic nervous system matures.

Small variations in heart rate and blood pressure have been recognised since the nineteenth century. Traube and Hering described fluctuations in blood pressure at the same frequency as respiration. Mayer described waves in blood pressure slower than those of respiration in animals with spontaneous respiratory movements. Guyton and Harris reported cycles of blood pressure at a frequency of 1 to 5/min in deteriorating animal preparations. Recently, Anthony et al described slow variability of cerebral blood flow velocity (CBFV) in both term and preterm newborn babies at a similar frequency. In short term recordings a highly variable CBFV has been implicated in the aetiology of intracranial pathology in the very low birthweight infant. Slow variability is important because in a recording of CBFV lasting only part of one minute its presence might lead to misinterpretation of beat to beat variability.

In adults and term newborn infants, slow variability is thought to be influenced by both thermoregulation and by baroreceptor reflex control mediated by the autonomic nervous system, and this may also be true for the preterm infant. Studies recording CBFV in adults over 30 minute periods do not report the presence of slow variability, suggesting that at some point of postconceptional age this becomes so damped as to be undetectable.

In this study we examined slow variability in CBFV in preterm infants in a longitudinal study extending to 14 weeks of postnatal age.

Patients and methods

PATIENTS

The study was carried out in the Special Care Baby Unit (SCBU) at the Rosie Maternity Hospital in Cambridge between December 1989 and October 1990. From a cohort of 74 consecutive admissions of babies with birth weight <1500 g, 30 remained longer than four weeks and were the subjects of this study. Infants were excluded only if they had a lethal congenital malformation, were admitted to the SCBU after 24 hours of age, or were less than the third centile for weight for gestational age. Informed consent was obtained from one or both parents. The study was approved by the district ethics committee.

METHODS

Recordings of CBFV were made using Duplex Doppler ultrasound (ATL Mk600) on day one and at weekly intervals thereafter until discharge. These were performed by a single operator. Doppler signals were processed using a microcomputer based system developed by Schindwein and Evans. This used a fast Fourier transform to calculate the peak velocity every 6.25 ms and passed the result to a second microcomputer (Apple Macintosh II running Labview) which was used to store the data. Each recording lasted one minute. Subsequently the velocity recording was visually inspected, and classified as showing variability at a frequency of between 1 and 5/min, showing no such variability, or as unclassifiable. Variability at frequencies above 5/min, probably due to the influence of respiration, was not considered in this study.

Amplitude of variability was then calculated by using the maximum systolic velocity present in the whole recording, subtracting the minimum systolic velocity, and dividing by the maximum systolic velocity. This was then expressed as a percentage.

We validated the visual classification of slow variability by comparing visual inspection with calculated values of maximum variability. A value of less than 10% maximum variability was used as a cut off for comparison with visual classification of no slow variability, as previous work has shown that short term variability is usually less than 10%.

Each week study babies also had standard...
views of cerebral ultrasound recorded by skilled independent operators who assessed them for periventricular haemorrhage and for ischaemic lesions (periventricular leucomalacia).

Results

Altogether 133 recordings were obtained from 30 babies of mean birth weight 1113 g (range 573–1486) and mean gestational age 27 weeks (range 24–31). Postnatal age extended from birth to a maximum of 14 weeks. Babies ranged from those never ventilated to those ventilated for more than 28 days (table 1).

Six infants suffered a periventricular haemorrhage of a grade greater than 2 (three grade 4, and three grade 3 according to the classification of Levene et al.14) Four infants had ischaemic lesions, one with huge multifocal cysts, two with single anterior cysts, and one with multiple anterior cysts. One required shunting for post-haemorrhagic hydrocephalus. Slow variability was usually obvious, with only four recordings out of the 133 being unclassifiable. There was good agreement between visual classification and calculated amplitude of less than 10% (table 2).

Slow variability was sometimes present for only part of the minute recorded. When more than one cycle was observed, the cycle length was not always the same, meaning that the variability was not occurring at a constant frequency. The occurrence of slow variability decreased with increasing postconceptional age. All babies showed slow variability up to 26 weeks' postconceptional age, whereas this was observed in only 70% at 32 weeks and at 37 weeks this had fallen to 25% (fig 1). The 95% confidence intervals for this trend showed a significant reduction with increasing postconceptional age, as seen in fig 2A; babies were divided into four groups of increasing postconceptional age. Each group contained approximately equal numbers of babies.

A similar trend was seen with respect to postnatal age; the proportion showing epochs of no slow variability increased with increasing postnatal age. When 95% confidence intervals were calculated there was a significant difference between groups 1 and 3 and between groups 2 and 4 (fig 2B). In this instance groups consisted of babies of increasing postnatal age divided into periods of four weeks.

There was also a significant trend for damping of the amplitude of slow variability with increasing postconceptional age and with increasing postnatal age. Regression analyses of decreasing amplitude of variability with increasing age showed a significant correlation coefficient for seven of the eight babies with more

![Graph showing presence of slow variability in CBFV related to postconceptional age.](http://adc.bmj.com/)

**Figure 1** Presence of slow variability in CBFV related to postconceptional age.

![Graph showing percentage of observations made in different postconceptional age groups.](http://adc.bmj.com/)

**Figure 2** Presence of slow variability in CBFV related to (A) postconceptional age and (B) postnatal age showing the percentage incidence and 95% confidence intervals.

<table>
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<tr>
<th>Table 1 Characteristics of study infants (n=30)</th>
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<tr>
<td><strong>Mean</strong></td>
</tr>
<tr>
<td>Birth weight (g)</td>
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<tr>
<td>Gestational age (weeks)</td>
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<tr>
<td>No of days ventilated</td>
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<tr>
<td>No of days in oxygen</td>
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<tr>
<td>Periventricular haemorrhage:</td>
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<tr>
<td>Grade 3</td>
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<tr>
<td>Grade 4</td>
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<td>Periventricular leucomalacia</td>
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<tr>
<th>Table 2 Comparison of visual inspection compared with measured amplitude of slow variation</th>
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<tr>
<td><strong>Validity of visual classification</strong></td>
</tr>
<tr>
<td>Maximum change in systemic velocity &lt;10%</td>
</tr>
<tr>
<td>(false positive)</td>
</tr>
<tr>
<td>Maximum change in systemic velocity &gt;10%</td>
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<tr>
<td>(true negative)</td>
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- Positive predictive value: 94.7%
- Negative predictive value: 93.3%
- Efficiency: 93.8%
- False positive: 1.9%
- False negative: 19.3%
than nine recordings. Figures 3 and 4 show the epochs of CBFV recorded from one baby over 11 weeks, together with an example of such a regression analysis.

The pattern of evolution did not seem to relate to brain injury in the small group who sustained it except that, of the three who developed periventricular leucomalacia, one was the only baby who never demonstrated slow variability; the others showed slow variability up to the time of discharge. In the six with periventricular haemorrhage no particular pattern could be seen.

The disappearance of slow variability in CBFV could not have been due to truncation of the peak systolic velocity; the automatic scaling within the software always allows the maximum velocity to be displayed.

Careful inspection reveals the appearance of slow variability in the diastolic velocity of cerebral blood flow. This was observed in 19 infants, generally after 31 weeks' postconceptional age.

Discussion

This longitudinal study confirms the presence of slow variability in the CBFV of babies in the frequency range 1.5 to 5 cycles/minute as observed by Anthony et al; it has also shown postnatal evolution of the activity.

Slow variability occurred in the majority of very low birthweight infants in the first weeks of life, being recorded at least once in 29 of the 30 babies. It appeared to be independent of the presence of respiratory distress syndrome and of severity of illness as some of the infants required no respiratory assistance while others required early inotropic support and ventilatory assistance for up to 44 days. As this study included only babies still in the nursery at 4 weeks of age it excluded early neonatal deaths. Also, although the study infants were entered from a consecutive cohort of babies of birth weight <1500 g, those still in the nursery by 2–3 months all had chronic lung disease so were a very selected group.

Perlman et al postulated that paralysis of sick preterm infants who had respiratory distress syndrome decreased beat to beat variability of CBFV and reduced the incidence of brain injury. In this study administration of pancuronium, fentanyl, or morphine in seven babies did not affect the presence of slow variability. The pattern of slow variability did not differ significantly between those without intracranial pathology and those with either periventricular haemorrhage or periventricular leucomalacia. The single baby who showed no slow variability had the most severe intracranial pathology of all babies studied. This would support the fact that slow variability is a normal phenomenon in newborn babies, absent only in cases of extreme pathology. Indeed absence of any variability in heart rate is an adverse prognostic indicator for respiratory distress syndrome. In severe respiratory distress syndrome, increased baseline heart rate and decreased variability may result from an increased concentration of circulating catecholamines which may represent abnormally high sympathetic activity.

Since the original description of slow variability in blood pressure at a frequency independent of the respiratory frequency by Mayer, two other areas of cyclical activity have been recognised in both heart rate and blood pressure: a peak around 0.05 Hz is thought to be related to fluctuations in peripheral vascular tone as-

Figure 3 Evolution of slow variability in baby F from 28–39 weeks' postconceptional age.

Figure 4 Regression analysis in baby F.
sociated with thermoregulation and that around 0·12 Hz to the frequency response of the baroreceptor reflex. Work in adults and in animals has shown that variability is influenced by thermoregulation and by pharmacological manipulation of the parasympathetic and sympathetic nervous systems. The renin-angiotensin system has also been shown to play a significant part in short term cardiovascular regulation, and blocking this system increases the amplitude of very slow cycles (0·04 Hz) in dogs.

High variability in the region of vasomotor influence (0·0–0·1 Hz) relative to that in the region of respiratory frequency (0·25–1·0 Hz) is suggested to be a risk factor for sudden infant death syndrome and may represent slow maturation of the autonomic nervous system, with relative over-representation of the parasympathetic element. In the study group of premature babies the reduction in amplitude of slow variability may represent a maturation of the balance between the two components of the autonomic nervous system. The cycle length of slow variability was variable, suggesting that several low frequency components were represented but the recording time was limited to 1 minute by the computer software. Much longer recordings would be required to resolve the frequency of several very slow variabilities with any degree of accuracy.

The significance of slow cycling in the diastolic velocity which appeared later in gestation in just over half of the babies has yet to be explained. We have no evidence that slow variability is harmful; further work may clarify the significance of this exaggerated normal response.

We would like to thank the nursing staff of the special care baby unit, Rosie Maternity Hospital, Cambridge, and Francis Griffiths for help with software.

1 Traube L. Uber periodische Tätigkeitsanderungen des vaso-

motorischen und Hemmings-Nervenenzemtres. Centraiblaut

2 Herin E. Uber den Einfluss der Atmung auf das Kreislauf.

3 Mayer S, Studien zur Physiologie des herzus und der

4 Gunton AC, Harris JW. Presororeceptor-autonomic oscillation:

5 Anthony MY, Evans DH, Leven MI. Cyclical variations in

6 Perlman J, McMenamin JB, Volpe JJ. Fluctuating cerebral

7 Hyndman BW, Kitney RI, Sayers BM. Spontaneous rhythms

8 Kitney RI. An analysis of the nonlinear behaviour of the

9 Schechtman VL, Harper RM, Klage KA. Development of

10 van Ravenswaai-Arts CMA, Hopman JCW, Kelle JAA,

11 Ellingsen I, Haage A, Niclasen M, Thoresen M, Wallac

12 Schindwein FS, Evans DH. A real-time autoregressive

13 Rennie JM, South M, Morley CJ. Cerebral blood flow

14 Levene M, Dubowitz LMS, De Crespy LCH. Classification

15 Perlman J, Goodman S, Kreusser KL, Volpe JJ. Reduction in

16 Jenkins JG, Reid MM, McClure BG. Study of heart rate

17 Cheek DB, Malinek M, Fruillon JM. Plasma adrenaline and

18 Akselrod S, Gordon D, Ubel FA, Shannon DG, Berger AC,

19 Kitney RI. New findings in the analysis of heart rate

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Arch Dis Child 1992 67: 412-415
doi: 10.1136/adc.67.4_Spec_No.412

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