Neonatal seizures: the Dublin Collaborative Study

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SUMMARY Asphyxial seizures occurred in 89 of 101 829 infants born alive at term (0·87/1000) in three large maternity hospitals from January 1980 to December 1984. These seizures were significantly associated with antenatal complications, primiparity, and prolonged pregnancy. Meconium staining of the amniotic fluid was also associated with asphyxial seizures, but there were high false positive (11%) and false negative (50%) rates. Fifteen of the infants who had seizures died (18%) and 21 (25%) were handicapped at 1 year. Outcome was most successfully predicted by the way the infant was feeding at 1–2 weeks. All infants taking more than half their estimated requirements by mouth at 1 week were normal, and those still being fed by tube at 2 weeks were handicapped.

Asphyxia is the commonest cause of brain injury in the perinatal period. Prevention, particularly in infants born at term, is the major goal of perinatal care. Perinatal asphyxia has been variously defined as including intrapartum events such as abruptio placentae, accidents with the cord, abnormal patterns of fetal heart rate, low fetal scalp pH, and the passing of meconium by the fetus. Definitions have also included assessments of the infant's poor condition at birth such as the need for and the duration of resuscitation, the Apgar score, the acid base measurement, and varying combinations of abnormal neurological signs including seizures.

Many of these indices correlate poorly with both proven intrapartum asphyxia (hypoxia, acidosis, and ischaemia) and with brain damage among survivors.1–4 A seizure occurring within 48 hours of birth in an infant born at term is the most rigorously validated event that reflects asphyxia that can be prevented by better obstetric technique. Such seizures are also the most strongly predictive of either neonatal death or survival with handicap.5,6 In this study, which was carried out in three large maternity hospitals, we attempted to document the incidence and outcome of neonatal asphyxial seizures with a view to predicting and therefore preventing them. By identifying reliable prognostic factors in the immediate neonatal period, we hoped to counsel parents more accurately.

Patients and methods

The study population comprised all infants born alive at more than 37 completed weeks' gestation in the Coombe, the National, and the Rotunda maternity hospitals during the five years January 1980 to December 1984. Seizure was diagnosed when the consultant paediatrician found evidence of one of the following: myoclonic, multifocal clonic, generalised tonic, or focal clonic seizures. Infants with subtle seizures or 'jitteriness' were excluded. All infants were observed daily for at least four days postpartum, and those causing concern were seen daily by a consultant. Infants suspected of having seizures were admitted to the neonatal intensive care unit for assessment including estimations of blood glucose, sodium, and calcium concentrations; full blood count; lumbar puncture, neurological examination, and (latterly) cranial ultrasound examination. Infants of 37 weeks' gestation or more whose perinatal history suggested seizure due to intrapartum asphyxia and who had a fit within 48 hours of delivery were included in the study. All three hospitals collect data prospectively on infants with asphyxial seizures, which are presented at a monthly perinatal morbidity conference. The next normally formed infant of the same gestational age born after each infant who had a seizure was used as the control. Data about outcome of survivors was obtained up to a minimum of one year from the records of the developmental follow up clinics at each hospital. A poor outcome among those who had seizures was defined as death, or handicap including cerebral palsy, mental retardation, epilepsy, or hearing or visual deficit.
Antenatal variables documented in index cases and controls included maternal age, height, weight, weight gain in pregnancy, parity, incidence of cigarette smoking, and specific obstetric complications (hypertension, proteinuria, and antepartum haemorrhage). Intrapartum variables analysed included spontaneous or induced onset of labour, oxytocin stimulation, meconium stained amniotic fluid, fetal heart rate (monitored either by intermittent auscultation or by continuous electronic monitoring), mode of delivery, (normal, instrumental, or caesarean), and gestational age.

Neonatal variables recorded were sex, birth weight, Apgar score, need for intubation or resuscitation, and interval to the onset of the first seizure. Management of infants with seizures was similar in all three hospitals and included treatment with phenobarbitone (10–20 mg/kg), phenytoin (10 mg/kg) and diazepam (0·1 mg/kg); prophylactic anticonvulsant drugs were not given.

In order to assess the prognostic significance of methods of feeding, infants were assigned to one of three groups (assuming a requirement of 150 ml/kg/day of milk): those taking at least half their requirements orally by 1 week of age; those taking at least half their requirements orally by 2 weeks of age, and those who were still receiving more than half the daily requirements by nasogastric tube at 2 weeks of age. The $\chi^2$ test was used to estimate the significance of differences in proportions.

Results

During the study period a total of 101 829 infants were born alive after more than 37 completed weeks' gestation; asphyxial seizures occurred within 48 hours of birth in 89 (0·87/1000). These were compared with 88 control babies. The antepartum variables that were significantly associated with the development of seizures are shown in table 1. There was no association between the development of seizures and either cigarette smoking or weight gain of less than 10 kg during pregnancy.

The intrapartum variables that were significantly associated with the development of seizures are shown in table 2. Meconium staining did not occur in half the babies with seizures though it was found in 11% of the controls and those who had seizures were more likely to be born by emergency caesarean section or with the assistance of forceps than control babies. There was no association between induction of labour and development of seizures.

The babies who had seizures weighed significantly less at birth than controls (table 2) but when the neonatal data of those who had seizures were analysed with regard to outcome (table 3), the association

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Comparison of antenatal details between the groups</th>
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<tbody>
<tr>
<td></td>
<td>Babies with seizures (n=89)</td>
</tr>
<tr>
<td>Maternal age:</td>
<td></td>
</tr>
<tr>
<td>&lt;20</td>
<td>8</td>
</tr>
<tr>
<td>20-24</td>
<td>21</td>
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<tr>
<td>25-29</td>
<td>27</td>
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<tr>
<td>30-34</td>
<td>16</td>
</tr>
<tr>
<td>&gt;35</td>
<td>17</td>
</tr>
<tr>
<td>Parity:</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>49</td>
</tr>
<tr>
<td>1-4</td>
<td>27</td>
</tr>
<tr>
<td>&gt;4</td>
<td>13</td>
</tr>
<tr>
<td>Complications</td>
<td>34</td>
</tr>
<tr>
<td>No complications</td>
<td>55</td>
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<thead>
<tr>
<th>Table 2</th>
<th>Comparison of intrapartum details and birth weight between the groups</th>
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<tbody>
<tr>
<td></td>
<td>Babies with seizures (n=89)</td>
</tr>
<tr>
<td>Amniotic fluid stained with meconium</td>
<td>44</td>
</tr>
<tr>
<td>Emergency caesarean section</td>
<td>24</td>
</tr>
<tr>
<td>Mid or high cavity forceps delivery</td>
<td>24</td>
</tr>
<tr>
<td>Gestation longer than 41 weeks</td>
<td>24</td>
</tr>
<tr>
<td>Birth weight (g):</td>
<td></td>
</tr>
<tr>
<td>&lt;3000</td>
<td>25</td>
</tr>
<tr>
<td>3000–3500</td>
<td>35</td>
</tr>
<tr>
<td>&gt;3500</td>
<td>29</td>
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<th>Table 3</th>
<th>Association between measurements and outcome in 89 babies who had seizures</th>
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<tr>
<td></td>
<td>Good outcome</td>
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<tr>
<td>Feeding (n=66):</td>
<td></td>
</tr>
<tr>
<td>No taking half feeds orally by 7 days</td>
<td>25</td>
</tr>
<tr>
<td>No taking half feeds orally between 7 and 14 days</td>
<td>22</td>
</tr>
<tr>
<td>No requiring tube feeding for &gt;14 days</td>
<td>0</td>
</tr>
<tr>
<td>No intubated at birth (n=50):</td>
<td></td>
</tr>
<tr>
<td>&lt;10 minutes</td>
<td>14</td>
</tr>
<tr>
<td>&gt;10 minutes</td>
<td>10</td>
</tr>
<tr>
<td>Apgar score at one minute (n=84):</td>
<td></td>
</tr>
<tr>
<td>≤5</td>
<td>34</td>
</tr>
<tr>
<td>&gt;5</td>
<td>14</td>
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<tr>
<td>Apgar score at five minutes (n=84):</td>
<td></td>
</tr>
<tr>
<td>≤5</td>
<td>9</td>
</tr>
<tr>
<td>&gt;5</td>
<td>39</td>
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between low birth weight and poor outcome did not reach significance. Not surprisingly low Apgar scores at one minute (p<0.05) and 5 minutes (p<0.001) and the need for intubation for longer than 10 minutes at birth (p<0.001), were significantly associated with a poor outcome, but the age of occurrence of seizures was not. In 66 infants the method of feeding at 1 and 2 weeks of age was documented; all infants who were satisfactorily feeding orally at 1 week were normal, and all those requiring tube feeding at day 14 were abnormal (p<0.001).

Of the 89 infants who had seizures 15 died (18%), 21 survived but with handicaps (24%), and five were lost to follow up (all five were in the group who had rapidly returned to normal within the first week, four of whom have also been normal at six weeks, one did not attend). About a third of the surviving infants were handicapped.

There were significant differences among the rates of seizures among the three hospitals (table 4), as well as in the proportion of infants of 42 weeks' gestation or more in the seizure group (0 to 60%). This may reflect differences in the populations but must also reflect different management policies. The suggestion finds some support in the significant differences among the three hospitals in the number of caesarean sections performed (p<0.001), the number of induced labours (p<0.001), and the number of forceps deliveries (p<0.001). Among the infants born at more than 41 weeks' gestation who developed seizures (n=24), seven (29%) had a poor outcome compared with 29 of 65 infants born at term (45%); this was not significant.

Discussion

Developments in perinatal medicine over the past decade have largely concerned the prevention of perinatal asphyxia and the more rigorous assessment of the association between preventable asphyxia and the incidence of long term brain damage—for example, cerebral palsy.3 4 Given that many of our assumptions regarding the causes of cerebral palsy, mental retardation, and epilepsy are based on outdated knowledge1 and that there are many different definitions of perinatal asphyxia in current use, it is not surprising that confusion exists about the perinatal prevention of handicap. In addition, the introduction and widespread use of continuous fetal heart rate monitoring in labour has been justified by claims that it halves perinatal 'morbidity'7 despite the fact that we have no reliable estimate of how many complications originate in the perinatal period nor how many are really preventable.

Of the various definitions of intrapartum asphyxia currently used including abnormalities of the fetal heart rate, low fetal scalp blood pH, and meconium staining of the amniotic fluid, not only are none strongly predictive of the condition of the baby at birth but their association with long term outcome is unknown.2 3 8 9 Of the various methods of defining 'poor condition at birth' (birth asphyxia), both a low Apgar score and the need for intubation (but not acid base measurement) have been shown to be only weakly predictive of long term handicap.10-14

The randomised controlled trial of intrapartum fetal heart rate monitoring carried out in Dublin5 was the first clinical study large enough to allow assessment of the validity of these various indices of perinatal asphyxia. This study strongly supported the hypothesis that neonatal seizures within 48 hours of delivery reflect intrapartum asphyxia. There were no significant differences between the monitored and unmonitored groups for abnormal neurological signs (other than seizures) in the neonatal period, low Apgar score, low pH in the umbilical vein, apnoea at birth, or endotracheal intubation. Dennis and Chalmers6 proposed that the early (within 48 hours) rate of seizures in infants born at term would provide a sensitive indicator of the quality of obstetric care, and the Dublin study of monitoring seems to support this. In addition, the fact that the incidence of seizures in the three hospitals in the present study were significantly different (0.55, 0.91, and 1.2/1000 babies born alive at term, respectively, p<0.05) supports the hypothesis that asphyxial seizures are at least partially preventable.

The rate of seizures in the present study of 0.87/1000 babies born alive at term (43% having a poor outcome) compares favourably with other similar studies. Levene et al reported a poor outcome (1/1000 live births at term14) among

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<th>Table 4</th>
<th>Incidence of seizures and obstetric interventions in the three hospitals January 1980–December 1984</th>
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<tr>
<td>Hospital A</td>
<td>Hospital B</td>
</tr>
<tr>
<td>Incidence of seizures/1000 live births</td>
<td>0.55</td>
</tr>
<tr>
<td>Percentage of induced labours</td>
<td>22.4</td>
</tr>
<tr>
<td>Percentage of forceps deliveries</td>
<td>6.8</td>
</tr>
<tr>
<td>Percentage of caesarean sections</td>
<td>7.8</td>
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</tbody>
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The p values refer to significant differences in group rates for each variable.
asphyxiated infants delivered in Leicester from 1980 to 1984, and rates of 'moderate' and 'severe'
postasphyxial encephalopathy of 1–15 and 1–0/1000 babies born alive at term, respectively. All the
former and many of the latter group would be expected to have neonatal seizures though some
would be 'subtle seizures'. Leveno et al15 reported a rate of seizures of 1/1000 live births weighing more
than 2500 g, and used this 'low' rate as one of the
main justifications of a rate of caesarean section of 18%.

The present study included five infants born at
term weighing less than 2500 g, and excluded a similar
number of infants who had seizures but weighed
more than 2500 g and were born prematurely.

MacDonald et al8 showed that 93% of infants of
more than 28 weeks' gestation who develop
asphyxial seizures do so within 48 hours of delivery.
That one hospital in this study had a rate of seizures of 0–55/1000 babies born alive at term with a rate of
caesarean section of 8% means that prevention of
intrapartum asphyxia does not depend on the
number of caesarean sections. This hospital had a
high rate of induction of labour (22%) directed at
prevention of prolonged pregnancy, and used con-
tinuous monitoring of the fetal heart rate in 33% of
labours in pregnancies at term. This raises the
question of whether continuous monitoring of the fetal
heart rate prevents asphyxial damage. In the trial of
intrapartum monitoring a significant reduction in
asphyxial seizures occurred in the group that was
monitored, though this effect was confined to the
15% of labours that lasted more than five hours.
Length of labour was not included in the present
study as it was felt that the retrospective assessment
of onset and duration of labour was insuffiently
accurate.

In infants born at term asphyxial seizures have
been shown to reflect preventable obstetrical intra-

partum asphyxia in a prospective randomised trial.5
The present study confirms the association between
asphyxial seizures and poor outcome (43%) and we
are now able to counsel parents on the likelihood of
handicap using the infant's neurological state and the
ability to feed normally as guides.

Asphyxial seizures should be identified as well as
asphyxial deaths (before and during labour, and in
the neonatal period) in all maternity units because
they are a sensitive indicator of the value, or
otherwise, of perinatal intervention.

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References
1 Freeman JM. Prenatal and perinatal factors associated with brain
2 Hey E. Foetal hypoxia and subsequent handicap: the problem of
establishing a causal link. In: Chamberlain GVP, Orr CJB, Sharp F, eds. Litigation and obstetrics and gynaecology.
3 Niswander K, Henson G, Elbourne D, et al. Adverse outcome
of pregnancy and the quality of obstetric care. Lancet
4 Paneth N, Stark RI. Cerebral palsy and mental retardation in
5 MacDonald D, Grant A, Sheridan-Pereira M, Boylan P,
Chalmers I. The Dublin randomized controlled trial of intrapar-
tum foetal heart rate monitoring. Am J Obstet Gynecol
6 Dennis J, Chalmers I. Very early neonatal seizure rate: a
possible epidemiological indicator of the quality of perinatal
7 Quilligan EJ, Paul RH. Foetal monitoring: is it worth it? Obstet
Gynecol 1975;45:96–100.
8 Taylor DJ, Howie PW, Davidson J, Davidson D, Drillien CM.
Do pregnancy complications contribute to neurodevelopmental
9 Dennis J. The long-term effects of intrapartum cerebral
damage. In: Crawford JW, ed. Risks of labour. Chichester: John
10 Nelson KB, Ellenberg JH. Apgar scores as predictors of chronic
12 Brann AW. Factors during neonatal life that influence brain
disorders. In: Freeman JM, ed. Prenatal and perinatal factors
associated with brain disorders. Bethesda: National Institutes of
13 Sykes GS, Molloy PM, Johnson P, et al. Do Apgar scores
14 Levene MI, Sands C, Grindulis H, Moore JR. Comparison of
two methods of predicting outcome in perinatal asphyxia.
15 Leveno KJ, Cunningham FG, Pritchard JA. Caesarean section:
an answer to the house of Horne. Am J Obstet Gynecol
1985;153:838–44.

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