Sleeping position and electrocortical activity in low birthweight infants

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Objective: To evaluate the effects of prone and supine sleeping positions on electrocortical activity during active (AS) and quiet (QS) sleep in low birthweight infants.

Design: Randomised/crossover study.

Setting: Infant Physiology Laboratory at Children’s Hospital of New York.

Patients: Sixty three healthy, growing, low birthweight (birth weight 795–1600 g) infants, 26–37 weeks gestational age.

Interventions: Six hour continuous two channel electrocortical recordings, together with minute by minute behavioural state assignment, were performed. The infants were randomly assigned to prone or supine position during the first three hours, and positions were reversed during the second three hours.

Outcome measures and results: Fast Fourier transforms of electroencephalograms (EEGs) were performed each minute and the total EEG power (TP), spectral edge frequency (SEF), absolute (AP) and relative (RP) powers in five frequency bands [0.01–1.0 Hz, 1–4 Hz, 4–8 Hz, 8–12 Hz, 12–24 Hz] were computed. Mean values for TP, SEF, AP, and RP in the five frequency bands in the prone and supine positions during AS and QS were then compared. In the prone sleeping position, during AS, infants showed significantly lower TP, decreased AP in frequency bands 0.01–1.0 Hz, 4–8 Hz, 8–12 Hz, 12–24 Hz, increased RP in 1–4 Hz, and a decrease in SEF. Similar trends were observed during QS, although they did not reach statistical significance.

Conclusions: The prone sleeping position promotes a shift in EEG activity towards slower frequencies. These changes in electrocortical activity may be related to mechanisms associated with decreased arousal in the prone position and, in turn, increased risk of sudden infant death syndrome.

Low birth weight (LBW) associated with premature delivery stands out as one of the most significant risk factors for SIDS.24 This suggests that systematic studies of effects of sleeping position and sleep states on electrocortical activity in this high risk group could provide new insights into the understanding of SIDS. The objective of this study was to determine the effects of prone and supine sleeping position on electrocortical activity during QS and AS in healthy, growing LBW infants. Our general hypothesis was that in the prone sleeping position, as compared with supine, LBW infants would exhibit patterns of electrocortical activity that would be characteristic of deeper sleep. More specifically we proposed that, in the prone position, low frequency (1–4 Hz) activity would be increased, high frequency (12–24 Hz) would be decreased, and that this shift in power would be associated with a lower spectral edge frequency (SEF; defined as the frequency below which 95% of the power in a spectrum is found).

METHODS

Study population

The study population consisted of 63 healthy, growing LBW infants (birth weight 795–1600 g; 26–37 weeks gestational age), all of whom were enrolled in a prospective, double blind, controlled study of effects of quality of dietary energy on rate and composition of weight gain. The studies were approved by the institutional review board, and written consent was obtained from parents of all infants. All infants were being maintained in room air, were free of apnoea of prematurity, and were receiving no cardiac or respiratory drugs. None had sonographic evidence of central nervous

Abbreviations: AP, absolute power; AS, active sleep; EEG, electroencephalograph; LBW, low birth weight; QS, quiet sleep; RP, relative power; SEF, spectral edge frequency; SIDS, sudden infant death syndrome; TP, total power

system pathology at the time of the studies. Table 1 shows the characteristics of the infants.

Experimental design
Infants were studied in the Infant Physiology Laboratory at Children’s Hospital of New York. Biweekly studies of about six hours duration were performed starting when the infant reached full enteral intake of 180 ml/kg/day. Each study comprised two sequential three hour periods of continuous recording of two-channel electrocortical activity. Simultaneously, behavioural assignment of sleep state was made for each minute of the study. Infants were randomly assigned to the supine or prone position for the first three hour epoch; the position was then reversed in the second three hour epoch. They remained in their assigned positions throughout the interfeeding period, and no further manipulations were performed.

Experimental protocol
Infants were brought to the laboratory at about 0730 at which time electrodes for recording electrocortical activity and vital signs were attached. They were then placed in a radiantly warmed, clear plastic incubator and maintained under thermoneutral conditions. No physical constraints such as swaddling were used. Studies began after the 0800 feed and continued until the 1400 feed. The studies were interrupted for the 1100 feed, after which sleeping position was changed. The volume and composition of the two feeds were identical.

Measurement of electrocortical activity
Six silver/silver chloride EEG electrodes were placed in a subset of the International 10–20 System montage. After the impedance of each electrode had been tested, the head was wrapped with a gauze headband to secure the electrodes. Two electrodes were placed in a subset of the International 10–20 System montage. After the six silver/silver chloride EEG electrodes were placed in a subset of the International 10–20 System montage. After the six silver/silver chloride EEG electrodes were placed in a subset of the International 10–20 System montage. After the six silver/silver chloride EEG electrodes were placed in a subset of the International 10–20 System montage. After the six silver/silver chloride EEG electrodes were placed in a subset of the International 10–20 System montage. After the six silver/silver chloride EEG electrodes were placed in a subset of the International 10–20 System montage. After the six silver/silver chloride EEG electrodes were placed in a subset of the International 10–20 System montage. After the six silver/silver chloride EEG electrodes were placed in a subset of the International 10–20 System montage. After the six silver/silver chloride EEG electrodes were placed in a subset of the International 10–20 System montage.

Assessment of sleep state
Behaviour codes were assigned each minute using a scoring system developed and validated in our laboratory. Briefly, AS was coded if one or more rapid eye movements were observed during the minute. In addition to small body movements typical of AS, movements of whole extremities and the torso were seen in this state. QS was designated when the infant was asleep without rapid eye movements and appeared “rag-doll” floppy and relaxed; movements were limited to startles and non-nutritive sucking or jaw jerks. Indeterminate state was coded when small body movements were observed, without rapid eye movements. Codes were also assigned for awake, crying, and feeding periods.

Table 1 Characteristics of study population (n = 63)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Mean (SD)</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gestational age (weeks)</td>
<td>30.5 (2.5)</td>
<td>26–37</td>
</tr>
<tr>
<td>Study age (weeks)</td>
<td>34.5 (1.5)</td>
<td>31–38</td>
</tr>
<tr>
<td>Birth weight (g)</td>
<td>1264 (232)</td>
<td>795–1600</td>
</tr>
<tr>
<td>Study weight (g)</td>
<td>1852 (156)</td>
<td>1349–2235</td>
</tr>
</tbody>
</table>

Values are mean (SD) (range).

DISCUSSION
The hypothesis guiding our study was that, in the prone position, EEG activity would be shifted in a direction that would be indicative of deeper sleep—that is, 1–4 Hz activity would be higher and 12–24 Hz activity would be lower in the prone versus the supine position. When infants were judged to be in QS, position had no significant effect on either AP or RP in any frequency band. In AS this hypothesis was supported. However, this interpretation requires examination of both AP and RP and consideration of the possible impact of...
movement artefact. As hypothesised, in AS, AP in the 12–24 Hz band was lower in the prone position. However, the expected increase in absolute 1–4 Hz activity was not found. We propose that this increase is due to interference from low frequency artefact associated with movements. This, in concert with reduced movement in the prone position, obscured position related differences in 1–4 Hz activity. The AP in 0.01–1 Hz nearly doubled in AS as compared with QS. This increase probably results from contributions of movement artefact to activity in low frequency bands. Consistent with this interpretation is the finding that absolute 1–4 Hz activity was not higher in QS as would be expected. However, when state differences in the effect of movement artefact were reduced by expressing power relative to TP, higher levels of 1–4 Hz activity in QS were apparent.

AP in the 0.01–1 Hz band was about 19% greater in the supine versus the prone position. Although we did not quantify body or eye movements during the study, it is our impression that infants move less in the prone position. Thus, as with state, effects of body position on absolute levels of low frequency activity may also be masked by movement artefact. However, dividing by TP (which is primarily due to low frequencies) resolved the expected position effect on 1–4 Hz activity. Decreases in TP, decreases in AP in higher frequencies, increases in RP in the low frequency band from a life threatening challenge encountered during sleep may play a role. Of particular interest is slow wave delta—that is, 1–4 Hz—EEG activity which is not only a prominent feature of QS but is present as early as 32 weeks gestation in preterm infants. The frequency distribution of the sleep EEG, such as increase in spectral power in the 12–15 Hz band during QS at 4–8 weeks postnatal age in siblings of SIDS victims as compared with 8–12 weeks in control infants. Schectmann and colleagues have shown increased integrated delta amplitude in the early morning hours in SIDS siblings relative to control infants at 3–4 months postnatal age, suggesting increased arousal thresholds in the morning in SIDS siblings. These abnormalities of sleep EEG patterns are of special interest, as most SIDS deaths occur during sleep periods, and a failure to arouse from a life threatening challenge encountered during sleep may play a role.

Whether sleeping position related changes in arousal are associated with changes in cortical function as reflected in EEG characteristics is not known. There is some evidence of accelerated appearance of several developmental characteristics of the sleep EEG, such as increase in spectral power in the 12–15 Hz band during QS at 4–8 weeks postnatal age in siblings of SIDS victims as compared with 8–12 weeks in control infants. Schectmann and colleagues have shown increased integrated delta amplitude in the early morning hours in SIDS siblings relative to control infants at 3–4 months postnatal age, suggesting increased arousal thresholds in the morning in SIDS siblings. These abnormalities of sleep EEG patterns are of special interest, as most SIDS deaths occur during sleep periods, and a failure to arouse from a life threatening challenge encountered during sleep may play a role.

### Table 2: Absolute electroencephalographic spectral power (µV²) in five frequency bands in prone and supine positions during quiet and active sleep

<table>
<thead>
<tr>
<th>Frequency band [Hz]</th>
<th>Quiet sleep</th>
<th>Active sleep</th>
<th>p Value</th>
<th>Quiet sleep</th>
<th>Active sleep</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.01–1</td>
<td>82.27 (36.37)</td>
<td>95.19 (59.77)</td>
<td>0.26</td>
<td>158.24 (58.04)</td>
<td>187.63 (75.94)</td>
<td>&lt;0.0003</td>
</tr>
<tr>
<td>1–4</td>
<td>14.30 (5.14)</td>
<td>14.99 (6.85)</td>
<td>0.28</td>
<td>17.72 (5.90)</td>
<td>16.30 (6.37)</td>
<td>0.08</td>
</tr>
<tr>
<td>4–8</td>
<td>1.67 (0.92)</td>
<td>1.85 (1.45)</td>
<td>0.34</td>
<td>1.68 (0.64)</td>
<td>1.90 (0.89)</td>
<td>&lt;0.0005</td>
</tr>
<tr>
<td>8–12</td>
<td>0.73 (0.38)</td>
<td>0.80 (0.51)</td>
<td>0.29</td>
<td>0.69 (0.28)</td>
<td>0.81 (0.43)</td>
<td>&lt;0.005</td>
</tr>
<tr>
<td>12–24</td>
<td>0.74 (0.37)</td>
<td>0.83 (0.54)</td>
<td>0.23</td>
<td>1.09 (0.44)</td>
<td>1.29 (0.67)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Values are mean (SD).

### Table 3: Relative electroencephalographic spectral power (%) in five frequency bands in prone and supine positions during quiet and active sleep

<table>
<thead>
<tr>
<th>Frequency band [Hz]</th>
<th>Quiet sleep</th>
<th>Active sleep</th>
<th>p Value</th>
<th>Quiet sleep</th>
<th>Active sleep</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.01–1</td>
<td>80.62 (6.08)</td>
<td>81.37 (6.10)</td>
<td>0.59</td>
<td>87.00 (3.56)</td>
<td>87.96 (3.55)</td>
<td>&lt;0.0003</td>
</tr>
<tr>
<td>1–4</td>
<td>15.32 (5.00)</td>
<td>14.68 (5.16)</td>
<td>0.58</td>
<td>10.27 (3.03)</td>
<td>9.24 (2.90)</td>
<td>&lt;0.0003</td>
</tr>
<tr>
<td>4–8</td>
<td>1.76 (0.73)</td>
<td>1.71 (0.71)</td>
<td>0.80</td>
<td>0.97 (0.32)</td>
<td>0.93 (0.31)</td>
<td>0.12</td>
</tr>
<tr>
<td>8–12</td>
<td>0.79 (0.35)</td>
<td>0.77 (0.32)</td>
<td>0.80</td>
<td>0.40 (0.14)</td>
<td>0.40 (0.16)</td>
<td>0.60</td>
</tr>
<tr>
<td>12–24</td>
<td>0.80 (0.33)</td>
<td>0.79 (0.32)</td>
<td>0.08</td>
<td>0.64 (0.23)</td>
<td>0.65 (0.28)</td>
<td>0.70</td>
</tr>
</tbody>
</table>

Values are mean (SD).

heart rate variability, increased orthosympathetic tone, decreased parasympathetic tone, and lower blood pressure. Thus the combination of impaired arousal and altered autonomic nervous system function are probably intertwined in such a way as to increase risk of death in this sleeping position.

Whether sleeping position related changes in arousal are associated with changes in cortical function as reflected in EEG characteristics is not known. There is some evidence of accelerated appearance of several developmental characteristics of the sleep EEG, such as increase in spectral power in the 12–15 Hz band during QS at 4–8 weeks postnatal age in siblings of SIDS victims as compared with 8–12 weeks in control infants. Schectmann and colleagues have shown increased integrated delta amplitude in the early morning hours in SIDS siblings relative to control infants at 3–4 months postnatal age, suggesting increased arousal thresholds in the morning in SIDS siblings. These abnormalities of sleep EEG patterns are of special interest, as most SIDS deaths occur during sleep periods, and a failure to arouse from a life threatening challenge encountered during sleep may play a role. Of particular interest is slow wave delta—that is, 1–4 Hz—EEG activity which is not only a prominent feature of QS but is present as early as 32 week gestational age and increases in amount and amplitude during QS throughout later gestation and early postnatal life. The presence of slow wave activity during sleep is indicative of deeper sleep (higher arousal thresholds) in adult humans and animals. It is likely that infants are also more resistant to arousal when their sleep is characterised by slow wave activity. Decreases in TP, decreases in AP in higher frequencies, increases in RP in the low frequency band (1–4 Hz), and a shift in the SEF to lower frequencies are major distinguishing features of QS versus AS. Prone sleeping position led to significant changes in all four of these variables in the direction of QS, suggesting that the prone position promotes a deeper AS and may be related to a higher arousal threshold. These findings are especially important, as LBW infants, of 30–39 weeks postconceptional age, spend 74% of their time in AS. There is limited information on spectral analysis of EEG activity in preterm infants. The frequency distribution of the
EEG activity and the SEF in our study infants was very similar to previously published observations of LBW infants. 23,24 Schramm and colleagues24 reported a reduction in the AU in various frequency ranges (0.4-19.5 Hz) during periods of central apnoeas in preterm and term infants. This reduction from baseline was present in the time periods before and after manifestation of an apnoea. The authors suggest that these lower voltages of electrical activity represent parallel deficits in arousal reaction. Similar to their findings, we observed lower spectral powers in the prone sleeping position during AS. These sleep related changes in power during prone positioning may result from vestibular mediated influences through the cerebellum to thalamic synchronizing means. 25 This process, in turn, may inhibit incoming messages and deprive the cerebral cortex of signals from the outside world and account for the impaired arousal response in the prone position. Any impair- ment in arousability or associated shift in the electrocortical activity towards that characteristic of QS when infants are in the prone position could contribute to the final pathway to SIDS. However, most would agree that an arousal deficit failure alone would not result in SIDS unless it was paired with a lethal challenge.

Many questions concerning the importance of arousal and SIDS remain unanswered. During arousal from sleep, heart rate, blood pressure, and ventilation are increased, and, importantly, a behavioural response is evoked, allowing movement away from a life threatening stimulus. Our data lend support to a "failure to arouse" hypothesis for SIDS in that LBW infants, sleeping in the prone position exhibit a shift in the electrocortical activity towards the slower frequencies. Further advances in this area could potentially contribute to further reduction in SIDS incidence and earlier detection of infants at highest risk.

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This study was performed in the Infant Physiology Laboratory at Children′s Hospital of New York, Columbia University, New York, NY.

Competing interests: none declared

REFERENCES


IMAGES IN NEONATAL MEDICINE

Prenatal three dimensional ultrasound detection of linear nevus sebaceous syndrome

A full term, male neonate was born to a 38 year old mother by uncomplicated vaginal delivery. Amnioacentesis had been performed, and a chromosome study revealed a normal male karyotype at 11 weeks gestation. A prenatal three dimensional sonogram showed two skin tag-like lesions over the left orbital region at 28 weeks gestation (fig 1).

Physical examination of the neonate found linear, well demarcated, yellowish, hairless plaques located on the forehead, scalp, and facial region. Two nodules over the lateral canthus and upper eyelid of the left eye resulted in ectropion (fig 2) There were no other cutaneous lesions. Magnetic resonance imaging of the brain showed hemimegalencephaly. The baby suffered a seizure attack about two weeks after birth.

Linear nevus sebaceous syndrome is a rare sporadic oculoneurocutaneous disorder, consisting of a spectrum of abnormalities involving the skin, central nervous system, eyes, and other systems. The major clinical manifestations include linear nevus sebaceous, seizure, and mental retardation. The aetiology of this disorder is still not identified. The major finding on brain image is hemimegalencephaly, which is characterised by congenital overgrowth of one cerebral hemisphere ipsilateral to the skin lesions. The affected brain has essentially no function, and is often associated with hemiparesis, early onset seizures, mental retardation, hemimacrocephaly, and severe encephalopathy clinically. Prenatal diagnosis is difficult but a fetal sonogram, especially a three dimensional image, may be useful for early detection of linear nevus sebaceous syndrome.

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Parental consent was obtained for publication of figure 2

Competing interests: none declared

REFERENCES