The Effect of Head Positioning and Head Tilting on the Incidence of Intraventricular Hemorrhage in Very Preterm Infants: A Systematic Review

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Keywords
Intraventricular hemorrhage · Head position · Head rotation · Premature infant · Preterm birth · Tilting

Abstract
Background: Despite advances in neonatal intensive care, germinal matrix-intraventricular hemorrhage (GMH-IVH) remains a frequent, serious complication of premature birth. Neutral head position and head tilting have been suggested to reduce the risk of GMH-IVH in preterm infants during the first 72 h of life. Objective: The aim of this study was to provide a systematic review of the effect of neutral head positioning and head tilting on the incidence of GMH-IVH in very preterm infants (gestational age ≤ 30 weeks). In addition, we reviewed their effect on cerebral hemodynamics and oxygenation. Methods: Literature was searched (June 2016) in the following electronic databases: CINAHL, Embase, Medline, SCOPUS, and several trial registers. Results: One underpowered trial studied the effect of head positioning on the incidence of GMH-IVH. This randomized controlled trial enrolled 48 preterm infants and found no effect on the occurrence of GMH-IVH. Three observational studies investigated the effect of head rotation and/or tilting on cerebral oxygenation in 68 preterm infants in total. Their results suggest that cerebral oxygenation is not significantly affected by changes in head positioning. The effect of head positioning and/or tilting on cerebral hemodynamics was described in 2 observational studies of 28 preterm infants and found no significant effect. Conclusions: There is insufficient evidence regarding the effect of head positioning and tilting on the incidence of GMH-IVH and cerebral hemodynamics and oxygenation in preterm infants. We recommend further research in this field, especially in extremely preterm and clinically unstable infants during the first postnatal days.

Introduction
Germinal matrix-intraventricular hemorrhage (GMH-IVH) is a major, frequently occurring complication of preterm birth. Of the extremely premature infants (gestational age [GA] <28 weeks), 20–25% will develop a GMH-IVH, with the incidence being inversely proportional to GA [1]. Typically, GMH-IVH originates from the germinal matrix, a highly vascularized collection of...
neuronal-glial precursor cells in the developing brain that involutes from about 26 weeks of gestation onwards [2]. The etiology of GMH-IVH in preterm infants is multifactorial. One key factor is the intrinsic fragility of the germinal matrix vasculature [2]. The delicate blood vessels easily rupture when rapid changes in cerebral perfusion occur. This may subsequently lead to bleeding into the ventricles (intraventricular hemorrhage). A second contributing factor is the vessel pattern of the venous system in this area. Due to the U-shaped alignment, the veins are prone to venous congestion, which can cause vessel damage and bleeding [3]. Thirdly, disturbances and fluctuations in cerebral blood flow (CBF) are common in preterm infants. Especially in sick or extremely preterm neonates, autoregulation of cerebral perfusion is impaired [4–7]. Preterm infants are thus less able to maintain a relatively constant blood flow to the brain when changes in cerebral perfusion pressure occur. Consequently, fluctuations in systemic blood pressure as well as postural changes could lead to alterations in CBF. Once a GMH-IVH has occurred, this may result in serious complications such as posthemorrhagic ventricular dilatation and periventricular hemorrhagic infarction. A large and/or complicated GMH-IVH is strongly associated with an adverse outcome, including disabilities and death [1, 3, 8–10]. Despite numerous efforts to prevent GMH-IVH in premature infants, the incidence of severe GMH-IVH has remained stable during the last few decades [11, 12].

Though seemingly harmless, routine caregiving events may affect cerebral perfusion and oxygenation in the preterm neonate [13]. In an attempt to avoid (rapid) fluctuations in CBF as well as intracranial pressure during routine care, several nursing interventions have been proposed. These nursing interventions are especially important during the first 72 h after birth, since GMH-IVH mostly develops during this time window [3, 14]. The first of these interventions consists of positioning the head of the infant in a neutral (i.e., midline) position, enabling optimal cerebral venous drainage through the internal jugular veins. The jugular veins are the major outflow paths for cranial blood. Head rotation to either side may lead to occlusion or obstruction of the jugular venous-drainage system at the ipsilateral side. Indeed, jugular phlebograms and catheterization studies in term infants and children have shown that rotating the head 90° to one side may result in torsion and complete compression of the internal jugular vein on the same side [15–17]. As a consequence, hampered venous drainage could lead to venous congestion, subsequent increase in intracranial pressure, altered cerebral oxygenation, and ultimately bleeding [18–21]. The second proposed intervention consists of elevating the head of the incubator 15–30° upwards (i.e., tilting) in order to facilitate venous outflow from the brain by promoting hydrostatic cerebral venous drainage [22]. A multidisciplinary focus group has identified maintaining a neutral head position together with 30° tilting as 1 of 10 potential practices for the prevention of GMH-IVH. The rationale behind this recommendation was the finding that the benchmark site with the lowest rate of GMH-IVH used this practice [23, 24].

**Objective**

Our aim was to provide a systematic review of studies assessing the influence of head positioning and tilting on the incidence of GMH-IVH, as well as on cerebral hemodynamics and cerebral oxygenation in preterm neonates. The latter two being important factors in the etiology of GMH-IVH. Near-infrared spectroscopy (NIRS)-monitored cerebral oxygenation, oxygen extraction, and cerebral hemodynamics are correlated to the risk of GMH-IVH [7, 25].

**Methods**

**Design**

This systematic review was conducted according to the guidelines of the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) [26].

**Criteria for Considering Studies for This Review**

**Types of Studies**

Randomized controlled trials (RCTs), quasi-RCTs, controlled clinical trials, as well as observational studies were eligible for inclusion. Reviews, poster presentations, conference papers, or single-case studies were excluded. Studies were eligible for inclusion if they reported an objective clinical outcome measure such as the incidence and severity of GMH-IVH, cerebral oxygenation, or cerebral hemodynamic parameters. Availability of the full text was imperative.

**Types of Study Population**

The target population of this review consisted of preterm neonates (GA ≤30 weeks).

**Types of Interventions**

Studies evaluating at least 1 of the following 2 interventions for preventing GMH-IVH were included:

1. Neutral head positioning
2. Head tilting
Types of Outcome Measures

Primary outcome:
1. Incidence of GMH-IVH diagnosed by cranial ultrasonography

Secondary outcomes:
1. Cerebral perfusion
2. Cerebral oxygenation

Data Collection and Analysis

Electronic Searches. The following electronic databases were searched: Medline, Embase, CINAHL, SCOPUS, Cochrane Central Register of Controlled Trials (CENTRAL), and ClinicalTrials.gov. The search strategies are outlined in Appendix A. To avoid missing studies, we did not use a search filter to differentiate between study types. There was neither restriction on the basis of publication date nor on publication status. We restricted our search to articles written in English, Dutch, German, and French. We performed the searches on June 6, 2016.

Searching Other Resources. To identify potential additional studies, reference lists from relevant reviews and papers were hand searched.

Selection of Studies. Titles and abstracts of the search results were independently assessed by 2 review authors (K.A.d.B.-M. and A.J.B.). Full copies of all potentially relevant studies were obtained. Decision on final inclusion after retrieval of full papers was made by 2 authors independently (K.A.d.B.-M. and A.J.B.). Disagreements were resolved by discussion with a 3rd review author (G.v.W.-M.).

Data Extraction. Two review authors extracted details of the included studies independently using a data extraction form. Any disagreements about data were resolved by consensus; if necessary a 3rd reviewer was consulted. The following data were extracted from each study: study design, setting, patient characteristics, data collection, results, conclusion, and quality assessment.

Quality Appraisal of Individual Studies. Two authors (K.A.d.B.-M. and A.J.B.) independently evaluated the methodological quality. Discrepancies in ratings were discussed between the reviewers until consensus was reached. The methodological quality of the RCT was rated according to the “Jadad scale” (Appendix B) and the risk of bias by the “Cochrane Collaboration tool” [27]. The Jadad scale is used to independently assess the methodological quality of an RCT, with emphasis on the quality of randomization and blinding. It consists of a questionnaire composed of three questions resulting in a score ranging from 0 to 5 points. The Cochrane Collaboration risk of bias tool assesses the risk of various forms of bias (e.g. selection bias, performance bias, attrition bias, and reporting bias). Each of the items can be classified as low-risk, unclear-risk, or high-risk bias. For the quality appraisal of observational studies the “Strengthening the Reporting of Observational Studies in Epidemiology (STROBE)” statement was used [28]. The STROBE checklist is detailed in Appendix C. Each item was rated as positive, plus/minus, or negative. To provide a final judgment about the article, scores were categorized. If less than 50% of the total score of 22 points was achieved an article was classified as “weak.” If an article achieved between 34 and 66% of the total score, it was classified as “moderate.” An article was classified as “strong” if the total score was higher than 66%.

Grading Quality of a Body of Evidence. The quality of evidence for each of the outcome parameters was evaluated using the GRADE approach. The GRADE system entails an assessment of the quality of a body of evidence for each individual outcome specifier four levels of quality. The highest quality rating is assigned to evidence from RCTs. However, the evidence from RCTs can be downgraded depending on factors. Observational studies are generally graded as “low quality”, but 3 factors may increase the quality level of a body of evidence (Appendix D) [29].

Results

Study Selection

The electronic database search yielded 864 articles. One additional article was identified through hand searching. The initial selection, based on title and abstract, included 29 records that seemed to fulfill the predefined criteria. After reading the full-text articles, 24 of these 29 articles did not meet the inclusion criteria and were therefore excluded (12 reports on children/infants with a GA >30 weeks, 7 described no relevant outcome parameters, 1 letter to the editor, 1 case report, and 3 poster abstracts). Finally, 5 studies were included in this systematic review [30–34]. Figure 1 provides a flowchart of the study selection.

Study Characteristics

The 5 included studies involved a total of 120 preterm infants undergoing positional interventions. The number of subjects ranged from 4 to 48 for each study. Three reports included only neonates with a GA <30 weeks [30, 33, 34]. Due to a wider variation in GA in 2 other studies, the range of GA at birth in all studies combined varied between 24 and 33 weeks. Two studies included infants with a postnatal age ≤72 h [30, 33]. Postnatal age in the other three studies varied considerably within, as well as between studies, ranging from 1 h to 3 weeks [31, 32, 34]. The designs of the included studies were predominantly prospective observational; there was 1 RCT. Table 1 gives an overview of the outcome parameters measured in the included studies. Table 2 provides an overview about the characteristics and results of the included studies.

Methodological Quality

The quality of the single RCT is presented in Table 2a. The Jadad score was 4/5, indicating adequate randomization and blinding. Evaluation of the quality of the RCT using the Cochrane Collaboration risk of bias tool indicated a low risk of bias. The results of the quality appraisal of the observational studies are presented in Table 3. The quality of the studies varied between moderate and strong. None of the studies fulfilled all 22 STROBE qual-

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Frequently observed weaknesses were a lack of power analysis and strategies to prevent any type of bias.

Study Description
The results of the included studies are described according to the primary and secondary outcome measures: incidence of GMH-IVH, cerebral oxygenation, and cerebral hemodynamics.

Incidence of GMH-IVH
Only 1 report studied the effect of head position (neutral versus 90° rotation) on the incidence of GMH-IVH. In this RCT, premature infants (mean GA 27 weeks) were...
<table>
<thead>
<tr>
<th>First author, year, country</th>
<th>Design, setting, parameter(s)</th>
<th>Participants</th>
<th>Results</th>
<th>Quality of the RCT/observational study</th>
<th>Classification of evidence according to GRADE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>a Regarding incidence GMH-IVH</strong>&lt;br&gt;Al-Abdi [30], 2011, Saudi Arabia</td>
<td>Pilot RCT NICU Ultrasound IVH incidence</td>
<td>n = 48 Mean GA 27 weeks (±1.3 SD) Intervention start: &lt;2 h after birth Duration: first 7 days of life 90% on mechanical ventilation</td>
<td>Incidence all IVH in midline (neutral) head position: 6/23 (26%) Incidence all IVH in lateral head position (90° rotation): 5/25 (20%) RR 1.30; 95% CI 0.46–3.70; p = 0.62 Incidence IVH grade III–IV Midline head position: 2/23 (9%) Lateral head position: 4/25 (12%) RR 1.40; 95% CI 0.61–3.37; p = 0.94</td>
<td>(1) Jadad score: 4/5 (2) Risk of bias Allocation concealment: ✓ Performance bias:✗ Attribution bias: ✓ Reporting bias: ✓</td>
<td>Moderate evidence (minus 1 point due to sparse data)</td>
</tr>
<tr>
<td><strong>b Regarding cerebral oxygenation</strong>&lt;br&gt;Ancora [31], 2010, Italy</td>
<td>Observational Cross-over NICU NIRS (TOI)</td>
<td>n = 24 Mean GA at birth: 27.5 weeks (±2.8 SD) Mean postnatal age at the time of the study: 10.3 days (± 7.9 SD) No mechanical ventilation</td>
<td>Horizontal, midline (neutral) head position: TOI 66.7% (SD 2.2) Horizontal lateral (90° rotation) head position: 5–10 min after rotation: TOI 68.1% (SD 2.1) No significant change Elevated (30°) midline (neutral) head position: 5–10 min after rotation: TOI 68.5% (SD 2.0) Elevated (30°) lateral (90° rotation): head position: TOI 67.2% (SD 1.9) No significant change</td>
<td>STROBE 16/22 (73%)</td>
<td>Low evidence (well-performed observational studies)</td>
</tr>
<tr>
<td>Elser [32], 2012, USA</td>
<td>Observational Cross-over NICU NIRS (rScO2)</td>
<td>n = 24 GA at the time of inclusion: &lt;32 weeks (mean and SD not given) Repeated measures (days 2, 5, 7, weekly thereafter) No information on respiratory support</td>
<td>Horizontal, head midline (neutral) position (101 observations): mean rScO2 72.9% (SD 9.6) Horizontal, head 45° rotated to the right (19 observations): 10–20 min after rotation: mean rScO2 76.0% (SD 8.5) No significant change Differences in cerebral oxygenation after positional change became smaller as postmenstrual age increased (data not specified)</td>
<td>STROBE 16/22 (73%)</td>
<td>Strong observational study</td>
</tr>
<tr>
<td>Liao [33], 2015, USA</td>
<td>Observational Cross-over NICU NIRS (rScO2)</td>
<td>n = 20 Mean GA at birth: 26.5 weeks (±1.7 SD) Mean postnatal age at the time of the study: 2 days (range: 1–3) 35% on mechanical ventilation</td>
<td>Elevated (30°) midline (neutral) head position: rScO2 72.7% (SD 6.4) Elevated (30°) 45° rotated left side head position: rScO2 71.7% (SD 4.8) (measured with sensor positioned on the left side of the head 10–20 min after rotation) p &lt; 0.05 No statistically significant differences measured on the right side of the brain and/or after rotation to the right side</td>
<td>STROBE 20/22 (91%)</td>
<td>Strong observational study</td>
</tr>
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</table>
randomized to maintain either a neutral (midline) head position or a lateral head position during the first 7 days of life. Their results were inconclusive due to the small sample size (underpowered) [30]. GMH-IVH developed in 26% (6/23) of the infants with a neutral head position versus 20% (5/25) of the infants with a lateral head position (risk ratio 1.30; 95% confidence interval 0.46–3.70; p = 0.62).

Cerebral Oxygenation
Three observational reports investigated the effect of head rotation on NIRS parameters. These 3 studies, including a total of 68 infants, collected the NIRS measurements at different time points following head rotation, ranging from 5 to 20 min after rotation. Ancora et al. [31] did not find any significant changes in the cerebral tissue oxygenation index (TOI) 5–10 min after head rotation from midline to the side. All infants were stable and none was mechanically ventilated. Elser et al. [32] found no statistically significant difference in cerebral regional oxygen saturation (rScO2) 10–20 min after head rotation. Their results did however indicate that rScO2 changes following head rotation were smaller with increasing GA (no quantitative data presented). In a study performed by Liao et al. [33], a small but statistically significant decrease in rScO2 of 1% was found 10–20 min after head rotation from the midline (neutral position) to the left side. Rotation to the right side did not result in a significant change [33]. The effect of tilting on cerebral oxygenation in premature infants was investigated in the study performed by Ancora et al. [31], who did not find a significant change in the cerebral TOI 5–10 min after 30° elevation of the head of the incubator.

Table 2 (continued)

<table>
<thead>
<tr>
<th>First author, year, country</th>
<th>Design, setting, parameter(s)</th>
<th>Participants</th>
<th>Results</th>
<th>Quality of the RCT/observational study</th>
<th>Classification of evidence according to GRADE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buckley [34], 2009, USA</td>
<td>Observational Cross-over NICU Diffuse correlation spectroscopy and Doppler ultrasound</td>
<td>n = 4 Mean GA at birth: 26.3 weeks (range: 25–27) Mean corrected GA at the time of the study: 29.2 weeks (range: 26–34) No information on respiratory status</td>
<td>In each infant 9 days of data sets No significant change in CBF during first 5 min after tilting (12° elevation) Small sample size</td>
<td>STROBE 12/22 (55%) Moderate observational study</td>
<td>(Very) low evidence (moderately performed observational study)</td>
</tr>
<tr>
<td>Ancora [31], 2010, Italy</td>
<td>Observational Cross-over NICU NIRS (nTHI)</td>
<td>n = 24 Mean GA at birth: 27.5 weeks (±2.8 SD) Mean postnatal age at the time of the study: 10.3 days (±7.9 SD) No mechanical ventilation</td>
<td>Horizontal midline (neutral) head position: nTHI 1.04 (SD 0.06) Horizontal lateral (90° rotation) head position: 5–10 min after rotation: nTHI 0.93 (SD 0.08) No significant change Elevated (30°) midline (neutral) head position: 5–10 min after rotation: nTHI 1.09 (SD 0.09) Elevated (30°) lateral (90° rotation) head position: nTHI 0.89 (SD 0.07) No significant change Subgroup analysis: in all newborns with GA ≤26 weeks nTHI was significantly higher in the midline (neutral) head position than the 90° rotated position (p &lt; 0.05) (data not specified)</td>
<td>STROBE 16/22 (73%) Strong observational study</td>
<td>Horizontal versus head tilting Horizontal midline (neutral) head position: nTHI 1.04 (SD 0.06) Elevated (30°) midline (neutral) head position: 5–10 min after elevation: nTHI 1.09 (SD 0.09) No significant change Horizontal lateral (90° rotation) head position: nTHI 0.93 (SD 0.08) Elevated (30°) lateral (90° rotation) head position: 5–10 min after elevation: nTHI 0.89 (SD 0.07) No significant change</td>
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</table>
Cerebral Hemodynamics

Two observational reports on a total of 28 infants investigated the effect of head position on parameters reflecting cerebral hemodynamics. First, Ancora et al. [31] did not find any significant changes in the cerebral normalized total hemoglobin index (nTHI) 5–10 min after head rotation from the midline to the side. A subgroup analysis was conducted in infants with a GA ≤ 26 weeks, which revealed that nTHI was significantly higher if the head was in the midline position as compared to 90° rotation. No quantitative data were presented on the number of neonates in this subgroup analysis or their clinical characteristics [31]. Their study also investigated the effect of tilting on cerebral hemodynamics and found no significant change in cerebral nTHI after 30° elevation of the head of the incubator [31]. Buckley et al. [34] repeatedly studied the effect of a small elevation of the head (12° tilting) in a small group of 4 infants. They did not find a significant alteration in microvascular blood flow (assessed by means of diffuse correlation spectroscopy) or macrovascular blood flow velocity in the middle cerebral artery (assessed by transcranial Doppler ultrasound).

Grading the Quality of a Body of Evidence

Incidence of GMH-IVH

The evidence regarding the effect of head rotation on the incidence of GMH-IVH was qualified as being “moderate”. Although the magnitude of the effect was investigated by a well-performed RCT, the sample size was considered too small and underpowered. Therefore, there is insufficient evidence that head rotation affects the incidence of GMH-IVH in premature infants with a GA ≤ 30 weeks. None of the studies investigated the effect of tilting on the incidence of GMH-IVH in premature infants. Therefore, there is insufficient evidence supporting a relationship between head elevation and the occurrence of GMH-IVH.

Cerebral Oxygenation

Three well-performed observational studies provided information about changes in cerebral oxygenation after head rotation [31–33]. One of these studies also provided information on the effect of tilting on cerebral oxygenation [31]. Combined the previous studies provide low quality of evidence that head rotation and/or head tilting in premature infants (GA ≤ 30 weeks) does not (importantly) affect cerebral oxygenation 5–20 min after rotation/elevation.

Cerebral Hemodynamics

Together the 2 observational studies represent (very) low quality of evidence that head rotation and/or tilting does not influence cerebral hemodynamics [31, 34].

Discussion

Our aim was to provide a systematic review of studies assessing the influence of maintaining a neutral head position and of head tilting on the incidence of GMH-IVH in very preterm neonates (GA ≤ 30 weeks). In addition, we reviewed the effect of these postural changes on cerebral hemodynamics and oxygenation in this subset of patients, since these factors are closely related to the development of GMH-IVH [7, 25]. We found moderate quality of evidence that the incidence of GMH-IVH is not influenced by maintaining a neutral head position in very preterm infants. Al Abdi et al. [30] conducted the first and only study investigating the effect of head position on the
occurrence of GMH-IVH in preterm infants with a GA <30 weeks during the first 7 days after birth. The results of this pilot study were inconclusive due to its small sample size.

Low-quality evidence indicated that there is no significant effect on cerebral oxygenation by head rotation and/or head tilting in preterm infants [31–33]. One study performed by Liao et al. [33] revealed a small statistically significant one-sided decrease in rScO₂ after head rotation to the left side. However, this decline of only 1% is unlikely to be of clinical significance.

Regarding cerebral hemodynamics, (very) low-quality of evidence showed no effect on cerebral hemodynamics after head rotation and/or head tilting [31, 34].

The studies included in this systematic review were heterogeneous, especially with regard to the characteristics of the participants (e.g., variation in respiratory support), type of intervention (e.g., degree of tilting and/or rotation), and type of data collected (e.g., NIRS or ultrasound). All studies were performed in small groups of clinically stable infants. Postnatal age of most infants was more than 1 week, which is important since autoregulation of cerebral perfusion improves with postnatal age, whereas the risk of developing a GMH-IVH declines after the first postnatal days.

None of the studies reported information on the occurrence of side effects of the postural changes such as respiratory distress or increased rate of apnea. Maintaining a neutral head position hampers a prone position. Since prone position is thought to facilitate breathing it is possible that maintaining a neutral head position might predispose infants to respiratory complications [35–37].

Implications for Clinical Practice

Presently, there is insufficient evidence regarding the effect of a neutral head position and/or head tilting on the incidence of GMH-IVH in preterm infants. We therefore can neither recommend nor refute the use of a neutral head position and/or head tilting in order to prevent GMH-IVH.

Implications for Research

Further research (preferably an RCT) is needed in larger groups of preterm infants, focusing on the effect of neutral head positioning and head tilting on the incidence of GMH-IVH as well as cerebral hemodynamics and oxygenation. Special attention should be given to unstable, ill, preterm infants during the first 72 h after birth, since these are at greatest risk of developing a GMH-IVH. In addition, GA may influence the vulnerability to the unfavorable effects of head rotation and/or horizontal head position [19, 22, 31, 32]. Therefore, extremely premature infants should be included in future research. Importantly, these future studies should include information on possible negative side effects such as increased work of breathing.

Limitations and Strengths

This review has several limitations. First, literature bias may be possible because only original, published studies were included. Results published in journals may differ systematically from those in reports, poster presentations, dissertations, or conference papers. Second, due to the lack of homogeneity among the study designs, types of measurement, and statistical analyses, it was not possible to pool the data for meta-analysis. Third, selection based on language could possibly have led to some degree of selection bias. Strengths of this review are the detailed transparent and structured data collection procedure according to the GRADE system. Second, there were no limitations set on publication date. Furthermore, two reviewers performed the selection process and the analysis of the methodological quality.

Conclusion

There is insufficient evidence regarding the effect of head positioning and/or tilting on the incidence of GMH-IVH and on cerebral oxygenation and/or hemodynamics in very preterm infants. Further research is recommended with special focus on the clinically unstable, extremely preterm infant during the first 72 h after birth.
### Appendix A

**Medline Search**


AND


AND


Restriction: newborn: birth – 1 month. Restriction: language: English, Dutch, German, and French

**Embase Search**

(prematurity/exp OR premature:ab,ti OR preterm:ab,ti OR neonatal:ab,ti OR neonate:ab,ti OR low birthweight/exp OR low birthweight:ab,ti OR infant:ab,ti OR newborn:ab,ti)

AND

("intracranial hemorrhage'/exp OR intracranial hemorrhage:ab,ti OR cerebral hemorrhage:ab,ti OR intraventricular hemorrhage:ab,ti OR 'brain hemorrhage'/exp OR brain hemorrhage:ab,ti OR periventricular hemorrhage:ab,ti OR subependymal hemorrhage:ab,ti OR intracranial bleeding:ab,ti OR intraventricular bleeding:ab,ti OR subependymal bleeding:ab,ti OR brain bleeding:ab,ti OR 'intracranial pressure'/exp OR intracranial pressure:ab,ti OR cerebral perfusion:ab,ti OR brain perfusion:ab,ti OR cerebral oxygenation:ab,ti OR cerebral saturation:ab,ti OR cerebral hemodynamics:ab,ti OR brain hemodynamics:ab,ti OR cerebral blood flow:ab,ti)

AND

(head position:ab,ti OR prone:ab,ti OR supine:ab,ti OR head rotation:ab,ti OR neck rotation:ab,ti OR head movement:ab,ti OR elevating:ab,ti OR elevation:ab,ti OR nursing:ab,ti OR handling:ab,ti OR positional:ab,ti OR posture:ab,ti OR postural:ab,ti OR tilting:ab,ti)

**CINAHL Search**

("infant, premature" (AB) OR premature (AB) OR preterm (AB) OR neonatal (AB) OR neonate (AB) OR “infant, low birth weight” (AB) OR low-birth-weight (AB) OR baby (AB))

AND

(intracranial hemorrhage (AB) OR cerebral hemorrhage (AB) OR intraventricular hemorrhage (AB) OR brain hemorrhage (AB) OR intraventricular bleeding (AB) OR intracranial pressure (AB) OR cerebral perfusion (AB) OR brain perfusion (AB) OR cerebral oxygenation (AB) OR cerebral saturation (AB) OR cerebral hemodynamics (AB) OR cerebral blood flow (AB))

AND

(head position (AB) OR prone (AB) OR supine (AB) OR head rotation (AB) OR neck rotation (AB) OR head movement (AB) OR handling (AB) OR positional (AB) OR posture (AB) OR postural (AB) OR tilting (AB) OR elevating (AB) OR nursing (AB))

**SCOPUS Search**

TITLE-ABS (prematurity) OR TITLE-ABS (premature) OR TITLE-ABS (preterm) OR TITLE-ABS (neonatal) OR TITLE-ABS (neonate) OR TITLE-ABS (low birth weight) OR TITLE-ABS (infant) OR TITLE-ABS (newborn)

AND

TITLE-ABS (intracranial hemorrhage) OR TITLE-ABS (cerebral hemorrhage) OR TITLE-ABS (intraventricular hemorrhage) OR TITLE-ABS (periventricular hemorrhage) OR TITLE-ABS (subependymal hemorrhage) OR TITLE-ABS (intracranial bleeding) OR TITLE-ABS (intraventricular bleeding) OR TITLE-ABS (subependymal bleeding) OR TITLE-ABS (brain bleeding) OR TITLE-ABS (intracranial pressure) OR TITLE-ABS (cerebral perfusion) OR TITLE-ABS (brain perfusion) OR TITLE-ABS (cerebral oxygenation) OR TITLE-ABS (cerebral saturation) OR TITLE-ABS (cerebral hemodynamics) OR TITLE-ABS (brain hemodynamics) OR TITLE-ABS (cerebral blood flow)

AND

TITLE-ABS (head position) OR TITLE-ABS (prone) OR TITLE-ABS (supine) OR TITLE-ABS (head rotation) OR TITLE-ABS (neck rotation) OR TITLE-ABS (head movement) OR TITLE-ABS (positional) OR TITLE-ABS (posture) OR TITLE-ABS (postural) OR TITLE-ABS (tilting)
Appendix B

Jadad Scale Scoring

Score: Assign a score of 1 point for each "yes" or 0 points for each "no"

<table>
<thead>
<tr>
<th>Question</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Question 1</td>
<td>Was the study described as randomized (this includes the use of words such as randomly, random, and randomization)?</td>
</tr>
<tr>
<td>Question 2</td>
<td>Was the study described as double-blind (blinding of patients and evaluators, not necessarily therapists)?</td>
</tr>
<tr>
<td>Question 3</td>
<td>Was there a description of withdrawals and dropouts (explicit statement that all included patients were analyzed or the number and reasons for dropouts in all groups are given separately)?</td>
</tr>
</tbody>
</table>

Give 1 additional point if:

| Question 1 | The method to generate the randomization sequence was described and appropriate (table of random numbers, computer generated) |
| Question 2 | The method of double blinding was described and appropriate (independent blinded assessors used, identical placebo or active placebo treatment, neither the person doing the assessment nor the study participant could identify the intervention being assessed) |

Deduct 1 point if:

| Question 1 | The method to generate the randomization sequence was described and inappropriate (e.g., alternate allocation to groups, according to date of birth, hospital number, etc.) |
| Question 2 | The method of double blinding was described and inappropriate (the person doing the assessment and/or the study participant could identify the intervention being assessed) |

Appendix C

STROBE

<table>
<thead>
<tr>
<th>Item No.</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Title and abstract</td>
<td>(a) Indicate the study's design with a commonly used term in the title or the abstract</td>
</tr>
<tr>
<td></td>
<td>(b) Provide in the abstract an informative and balanced summary of what was done and what was found</td>
</tr>
<tr>
<td>Introduction</td>
<td>Explain the scientific background and rationale for the investigation being reported</td>
</tr>
<tr>
<td>Objectives</td>
<td>State specific objectives, including any prespecified hypotheses</td>
</tr>
<tr>
<td>Methods</td>
<td>Present key elements of study design early in the paper</td>
</tr>
<tr>
<td>Setting</td>
<td>Describe the setting, locations, and relevant dates, including periods of recruitment, exposure, follow-up, and data collection</td>
</tr>
<tr>
<td>Participants</td>
<td>(a) Cohort study: give the eligibility criteria, and the sources and methods of selection of participants; describe methods of follow-up</td>
</tr>
<tr>
<td></td>
<td>Case-control study: give the eligibility criteria, and the sources and methods of case ascertainment and control selection; give the rationale for the choice of cases and controls</td>
</tr>
<tr>
<td></td>
<td>Cross-sectional study: give the eligibility criteria, and the sources and methods of selection of participants</td>
</tr>
<tr>
<td></td>
<td>(b) Cohort study: for matched studies, give matching criteria and number of those exposed and unexposed</td>
</tr>
<tr>
<td></td>
<td>Case-control study: for matched studies, give matching criteria and the number of controls per case</td>
</tr>
<tr>
<td>Variables</td>
<td>Clearly define all outcomes, exposures, predictors, potential confounders, and effect modifiers; give diagnostic criteria, if applicable</td>
</tr>
<tr>
<td>Data source/measurement</td>
<td>For each variable of interest, give sources of data and details of assessment (measurement); describe comparability of assessment methods if there is more than 1 group</td>
</tr>
<tr>
<td>Bias</td>
<td>Describe any efforts to address potential sources of bias</td>
</tr>
<tr>
<td>Study size</td>
<td>Explain how the study size was arrived at</td>
</tr>
</tbody>
</table>
Item No. | Recommendation
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Quantitative variables 11 | Explain how quantitative variables were handled in the analyses. If applicable, describe which groupings were chosen and why

Statistical methods 12 | (a) Describe all statistical methods, including those used to control for confounding
(b) Describe any methods used to examine subgroups and interactions
(c) Explain how missing data were addressed
(d) **Cohort study:** if applicable, explain how loss to follow-up was addressed
**Case-control study:** if applicable, explain how matching of cases and controls was addressed
**Cross-sectional study:** if applicable, describe analytical methods taking account of the sampling strategy
(e) Describe any sensitivity analyses

Results Participants 13 | (a) Report numbers of individuals at each stage of the study, e.g., numbers potentially eligible, examined for eligibility, confirmed eligible, included in the study, completing follow-up, and analyzed
(b) Give reasons for nonparticipation at each stage
(c) Consider use of a flow diagram

Descriptive data 14 | (a) Give characteristics of study participants (e.g., demographic, clinical, and social) and information on exposures and potential confounders
(b) Indicate number of participants with missing data for each variable of interest
(c) **Cohort study:** summarize follow-up time (e.g., average and total amount)

Outcome data 15 | **Cohort study:** report numbers of outcome events or summary measures over time
**Case-control study:** report numbers in each exposure category, or summary measures of exposure
**Cross-sectional study:** report numbers of outcome events or summary measures

Main results 16 | (a) Give unadjusted estimates and, if applicable, confounder-adjusted estimates and their precision (e.g., 95% CI); make clear which confounders were adjusted for and why they were included
(b) Report category boundaries when continuous variables were categorized
(c) If relevant, consider translating estimates of relative risk into absolute risk for a meaningful time period

Other analyses 17 | Report other analyses done – e.g., analyses of subgroups and interactions, and sensitivity analyses

Discussion Key results 18 | Summarize key results with reference to study objectives

Limitations 19 | Discuss limitations of the study (taking into account sources of potential bias or imprecision); discuss and both direction and magnitude of any potential bias

Interpretation 20 | Give a cautious overall interpretation of results considering objectives, limitations, multiplicity of analyses, results from similar studies, and other relevant evidence

Generalizability 21 | Discuss the generalizability (external validity) of the study results

Other information Funding 22 | Give the source of funding and the role of the funders for the present study and, if applicable, for the original study on which the present article is based

An Explanation and Elaboration article discusses each checklist item and gives methodological background and published examples of transparent reporting (freely available on the Web sites of PLoS Medicine [www.plosmedicine.org], Annals of Internal Medicine [www.annals.org], and Epidemiology [www.epidem.com]). Information on the STROBE Initiative is available at www.strobe-statement.org.

* Give information separately for cases and controls in case-control studies and, if applicable, for exposed and unexposed groups in cohort and cross-sectional studies.
### Appendix D

#### Grade System

<table>
<thead>
<tr>
<th>Levels of quality of body of evidence</th>
<th>Quality rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>RCTs or double-upgraded observational studies</td>
<td>High</td>
</tr>
<tr>
<td>Downgraded RCTs or upgraded observational studies</td>
<td>Moderate</td>
</tr>
<tr>
<td>Double-downgraded RCTs or observational studies</td>
<td>Low</td>
</tr>
<tr>
<td>Triple-downgraded RCTs, downgraded observational studies, or case series/reports</td>
<td>Very low</td>
</tr>
</tbody>
</table>

**Factors that may decrease the quality level of a body of evidence**

1. Limitations in the design and implementation of available studies suggesting a high likelihood of bias (minus one or two levels depending on the degree of limitations)
2. Indirectness of evidence (minus one or two levels depending on the degree of uncertainty about directness)
3. Unexplained heterogeneity or inconsistency of results (minus one level)
4. Imprecision of results (wide confidence intervals) or sparse data (minus one level)
5. High probability of publication bias (minus one level)

**Factors that may increase the quality level of a body of evidence**

1. Large magnitude of effect (plus one level)
2. All plausible confounders would have reduced a demonstrated effect or suggest a spurious effect when results show no effect (plus one level)
3. Dose-response gradient (plus one level)

**Grade of evidence definition**

<table>
<thead>
<tr>
<th>Grade</th>
<th>Definition</th>
</tr>
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<tbody>
<tr>
<td>High</td>
<td>Further research is unlikely to change the confidence in the estimate of the effect</td>
</tr>
<tr>
<td>Moderate</td>
<td>Further research is likely to have an important impact on the confidence in the estimate of the effect and may change the estimate</td>
</tr>
<tr>
<td>Low</td>
<td>Further research is very likely to have an important impact on the confidence in the estimate of the effect and is likely to change the estimate</td>
</tr>
<tr>
<td>Very low</td>
<td>Any estimate of effect is very uncertain</td>
</tr>
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</table>

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**References**

Head Position and Prevention of Neonatal Intraventricular Hemorrhage

Neonatology 2017;111:267–279
DOI: 10.1159/000449240