Head Ultrasound (HUS) Screening in Premature Infants

PEDIATRIC NEWBORN MEDICINE CLINICAL PRACTICE GUIDE LINE
Clinical Practice Guideline: Head Ultrasound (HUS) Screening in Premature Infants

Points of emphasis/Primary changes in practice:
1. Clear indications for HUS screening in asymptomatic premature infants
2. Identification of the timing and frequency of HUS screening
3. Expanding the measurements performed on HUS and providing recent reference values.

Rationale for change:
Although HUS is routinely used in NICU, there has not been an updated guideline to minimize variability in practice.

Questions? Please contact: The Neurocritical Care Working Group
This is a clinical practice guideline. While the guideline is useful in approaching the use of HUS screening in premature infants, clinical judgment and/or new evidence may favor an alternative plan of care, the rationale for which should be documented in the medical record.

Introduction

The incidence of Germinal Matrix- Intraventricular Hemorrhage (GM-IVH) in very low birth weight infants (< 1500 grams at birth) is about 25% with the incidence of the severest forms of IVH (grades III and IV) being approximately 5%. About 50% of infants with GM-IVH are “clinically silent” and show no abnormal signs indicative of the lesion. 1-3 About 30–50% of infants with a large IVH develop post-hemorrhagic ventricular dilatation (PHVD) and around 20–40% of infants with a severe GMH–IVH will consequently need a permanent ventriculoperitoneal shunt. 4,5 On the other hand, periventricular leukomalacia seems to be the most important determinant of neurologic morbidity seen in preterm infants who survive the neonatal period. Although the cystic form, with focal necrotic lesions, is usually associated with the development of cerebral palsy (CP) in infancy, the more diffuse form, which is not well visualized with cranial ultrasound, may relate to cognitive and behavioral problems.6 Finally, in more recent years, cerebellar hemorrhage has been recently recognized as an important complication of extreme prematurity and is associated with increased morbidity and mortality.7,8

Head ultrasound via the cranial fontanels allows for rapid bedside evaluation of the neonatal brain. The standard views for HUS are the sagittal and coronal views through the anterior fontanel. In addition, the mastoid fontanel allows better visualization of the posterior fossa and brainstem, 9,10 while the posterior fontanel allows for better visualization of the trigone and occipital horn of the lateral ventricles as well as the posterior fossa.11 Major abnormalities on HUS (including grade III to IV IVH, or cystic PVL) are predictive of cerebral palsy and neurodevelopmental impairments.12-19

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Role of HUS screening in Premature Infants

1- Germinal Matrix- Intraventricular Hemorrhage (GM-IVH)

The classic grading system of GM-IVH was initially described by Papile et al. 20 Findings were graded on a scale from I to IV. Periventricular hemorrhagic infarction is the term that substituted “grade IV IVH” as it represents a distinct pathological entity caused by obstruction of blood flow in the terminal veins leading to venous infarction and hemorrhage into the surrounding tissue. 21,22 The onset of GM-IVH on HUS is by the first day of life in at least 50% of affected infants, with up to 90% of lesions present by 72 hours 23,24 and all lesions visible by 7 days. 12 However, further progression of an initial IVH occurs in approximately 20% to 40% of the affected infants, with the maximal extent generally seen by 3 to 5 days after the initial diagnosis. 25,26

2- Ventriculomegaly

Ventriculomegaly is detected by HUS with excellent sensitivity. 27 Levene described a ventricular index that is defined as the distance in millimeters between the midline and lateral border of the smaller lateral ventricle in the coronal plane at the level of the foramen of Monro. 28 Ventricular diameter > 4 mm above the 97th percentile is considered severe and has been used as an inclusion criterion for different studies as a marker of progressive ventricular dilatation requiring intervention. 29-31 An updated graph with more extremely immature infants from 23 weeks GA was recently published. 32 Davies et al. 33 proposed a more detailed methodology that includes measuring different distances including anterior horn width (AHW), thalamo-occipital distance (TOD), third ventricle width and fourth ventricle width and length. A newer reference range was also published in 2012. 34

3- White Matter Injury (Periventricular Leukomalacia)(PVL)

White matter injury appears on HUS as increased periventricular echogenicity that may or may not develop into cysts. Cyst development (c-PVL) is often not detectable by HUS until 4 to 6 weeks following injury as coagulation necrosis and cellular loss evolve over time. It is to be noted that the incidence of c-PVL has declined significantly over the years. 35 White matter injury is better identified and evaluated using MRI. Term equivalent MRI is indicated for premature infants at highest risk for brain injury. 36
4- Cerebellar Hemorrhage

The sensitivity and specificity of HUS to detect cerebellar hemorrhage is variable because the cerebellum is echogenic and often difficult to distinguish from infarctions and hemorrhages. With careful attention to asymmetry in echogenicity and the use of mastoid or posterior fontanelle views, these lesions are more easily visualized.\textsuperscript{37,38}

Imaging Protocols

There is no universally accepted protocol for HUS screening in the preterm infant. The Report of the Quality Standards Subcommittee of the American Academy of Neurology and the Practice Committee of the Child Neurology Society states: “Routine screening cranial ultrasonography should be performed on all infants of 30 weeks’ gestation once between 7 and 14 days of age and should be optimally repeated between 36 and 40 weeks’ postmenstrual age”.\textsuperscript{39} In Europe, HUS may be done more frequently. Some centers perform HUS as soon as possible after admission, at least once a week until discharge, and again at term equivalent during the first follow up visit.\textsuperscript{12} This variation in frequency and timing of imaging may affect the ability of HUS to detect abnormality and subsequently predict outcome.

Interpretation of Normal Head Ultrasound

While head ultrasound is useful in detecting hemorrhage, ventricular dilatation and cystic changes, a “normal head ultrasound” does not exclude brain injury and does not imply normal outcome. In a report by NICHD, about one third of Extremely Low Birth Weight Infants with normal head ultrasounds had either significant mental developmental delay or cerebral palsy at the age of 18 to 22 months’ corrected age.\textsuperscript{40} This limitation should be acknowledged in documentation and in parents counselling.
The following guidelines were based on the consensus on the importance of HUS screening for premature infants at highest risk of brain injury. These guidelines are aimed at those with no clinical signs or symptoms. In presence of clinical signs or symptoms, suggestive of intracranial pathology, the clinician will request HUS regardless of screening criteria.

1. Purpose
To provide guidelines for Head Ultrasound (HUS) Screening in premature infants admitted to BWH NICU.

2. Indications of HUS screening:
   1- Gestational Age < 32 weeks, or
   2- Birth Weight < 1500 gm

3. Timing of Screening HUS:
   a- Less than 28 weeks GA
      1- At 1 day of life
      2- At 3 days of life
      3- At 1 week of life
      4- At 1 month of age
      5- At 36 weeks’ Postmenstrual age (if no term equivalent MRI completed)
   b- From 28 weeks to less than 32 weeks GA
      1- At 3 days of life
      2- At 1 week of life
      3- At 1 month of age
      4- At 36 weeks’ Postmenstrual age (if no term equivalent MRI completed)

4. Frequency of HUS if abnormal:
HUS will be repeated twice a week until abnormality is stable or resolving then routine US screening will be continued as scheduled.
However, in case of ventricular dilatation up to 3 HUS/week in the first 3 week of life will be considered.

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5. **Technical Guidelines:**

The following imaging protocol will be followed:

- **Still and Video Clips Views:**
  - Anterior fontanelle
    - Coronal
    - Sagittal
    - Parasagittal R and L
  - Posterior fontanelle view
  - Mastoid view

6. **Measurements:**

In case of presence of intraventricular hemorrhage (IVH) the following measurements will be completed and reported as measurement in mm:

1. Ventricular Index (VI) (R and L)
2. Anterior Horn Width (AHW) (R and L)
3. Thalamo-occipital Distance (TOD) (R and L)

7. **Appendix:**

**Summary of the Significance of Lateral Ventricles Measurement on Head Ultrasound**
Appendix 1: Ventricular Index (VI)

**Definition:** The distance in millimeters between the midline and the lateral border of the anterior horn of lateral ventricles. It is measured in the coronal view at the level of foramen of Monro. (Figure 1A)

![Image of ventricular index](image1a.png)

**Significance:**

**Normative Values:** Normative values has been established in 1981 using cross-sectional data from infants from 27-42 weeks GA. An updated graph with more extremely immature infants from 23 weeks GA was recently published 32 (Figure 1B)

![Image of ventricular width centile chart](image1b.png)

**Abnormal VI:** VI > 4 mm above 97th percentile was reported to be associated with about 50–60% being shunt dependent, over 60% disabled, and around 20% mortality. 41 For that reason, VI > 4 mm above 97th percentile was used as intervention criteria for many trials 29,31,42

However, in a retrospective study, early intervention (VI > 97th percentile) was associated with 16% need for VP shunt compared to 62% if late intervention (VI > 4 mm above 97th percentile) criteria was used 43

**In the ongoing ELVIS (early versus late ventricular intervention study) study:**
- Criteria for early intervention are: VI > p97 + AHW > 6 mm and/or TOD > 24 mm
- Criteria for late intervention are: VI > p97 + 4 mm and AHW > 10 mm

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Appendix 2: Anterior Horn Width (AHW)

**Definition:** The distance in millimeters between the medial wall and floor of the lateral ventricle. It is measured at the widest point of the anterior horn of the lateral ventricles, in the coronal view at the level of foramen of Monro. (Figure 2A and 2B) 

**Significance:** Rounding “ballooning” of the frontal horns and an increase in AHW could be an early sign of increased ICP. While several authors reported no or just minimal change in AHW with GA, an evident increase in AHW was noted by others.

**Normative Values:** Figure 2C can be used in first HUS while figure 2D can be used for follow up. 

- **AWH of 5 or less is acceptable.** In the majority of neonates, the AHW is less than 3 mm. An AHW between 3 and 5 mm was not associated with neurodevelopmental impairment at a single follow up visit in a small group of 13 infants within the first year of life.

- **Abnormal AHW:** Values exceeding 6 mm, however, are associated with ventricular ballooning and might suggest the need for treatment.

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Appendix 3: Thalamo-occipital Distance (TOD)

**Definition:** is the distance in mm from the outermost point of the thalamus, at its junction with the choroid plexus, to the outermost part of the occipital horn posteriorly in the oblique parasagittal view. (Figure 3A and 3B) \(^{44}\)

![Image of TOD measurement](image)

**Significance:**
Enlargement of Occipital horn is usually could be detected before is usually more significant than anterior horn dilatation. Sometimes, it may represent the only site of ventricular dilatation. Whether it is stable across GA has been controversial \(^{45}\)

**Normative Values:** Figure 3C can be used in first HUS while figure 3D can be used for follow up. \(^{34}\)
According to Davies et al., the mean is 16.7 ± 4 mm (Range 8.7-24.7) \(^{44}\)

**Abnormal TOD:** Based on multiple references a **value ≥ 25 mm** is considered abnormal
References:


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