Fetal Intraventricular Hemorrhage in Open Neural Tube Defects: Prenatal Imaging Evaluation and Perinatal Outcomes

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ABSTRACT

BACKGROUND AND PURPOSE: Fetal imaging is crucial in the evaluation of open neural tube defects. The identification of intraventricular hemorrhage prenatally has unclear clinical implications. We aimed to explore fetal imaging findings in open neural tube defects and evaluate associations between intraventricular hemorrhage with prenatal and postnatal hindbrain herniation, postnatal intraventricular hemorrhage, and ventricular shunt placement.

MATERIALS AND METHODS: After institutional review board approval, open neural tube defect cases evaluated by prenatal sonography between January 1, 2013 and April 24, 2018 were enrolled (n = 504). The presence of intraventricular hemorrhage and gray matter heterotopia by both prenatal sonography and MR imaging studies was used for classification. Cases of intraventricular hemorrhage had intraventricular hemorrhage without gray matter heterotopia (n = 33) and controls had neither intraventricular hemorrhage nor gray matter heterotopia (n = 229). A total of 135 subjects with findings of gray matter heterotopia were excluded. Outcomes were compared with regression analyses.

RESULTS: Prenatal and postnatal hindbrain herniation and postnatal intraventricular hemorrhage were more frequent in cases of prenatal intraventricular hemorrhage compared with controls (97% versus 79%, 50% versus 25%, and 63% versus 12%, respectively). Increased third ventricular diameter, specifically >1mm, predicted hindbrain herniation (OR = 3.7 [95% CI, 1.5–11]) independent of lateral ventricular size and prenatal intraventricular hemorrhage. Fetal closure (n = 86) was independently protective against postnatal hindbrain herniation (OR = 0.04 [95% CI, 0.01–0.15]) and postnatal intraventricular hemorrhage (OR = 0.2 [95% CI, 0.02–0.98]). Prenatal intraventricular hemorrhage was not associated with ventricular shunt placement.

CONCLUSIONS: Intraventricular hemorrhage is relatively common in the prenatal evaluation of open neural tube defects. Hindbrain herniation is more common in cases of intraventricular hemorrhage, but in association with increased third ventricular size. Fetal closure reverses hindbrain herniation and decreases the rate of intraventricular hemorrhage postnatally, regardless of the presence of prenatal intraventricular hemorrhage.

ABBREVIATIONS: NTD = neural tube defects; BPD = biparietal diameter; GA = gestational age by sonography; IVH = intraventricular hemorrhage; HC = head circumference; HH = hindbrain herniation; GMH = gray matter heterotopia; ONTD = open neural tube defects; MMC = myelomeningocele; MS = me-}

0neural tube defects (NTD) can be classified as open (eg, myelomeningocele [MMC] and myeloschisis [MS]) or closed, and detailed radiologic evaluation is key to their differentiation. Fetal sonography is effective in the assessment and characterization of NTD, with important implications for patient counseling and management. Fetal MR imaging provides supplementary or additional diagnostic information in these disorders. Both imaging modalities have similar performance in the evaluation of NTD prenatally and when compared with postnatal imaging. The association between open neural tube defects (ONTD) and hindbrain herniation (HH) has long been described. In animal studies, surgically created MMC produced HH that could be reversed with prenatal repair. The Management of Myelomeningocele Study (MOMS) trial demonstrated clinical
effectiveness and further studies have supported these initial results. In the MOMS trial, the safety and efficacy of fetal closure was proved with slight outcome discrepancies in subsequent studies. Prenatal identification of HH is crucial for patient counseling and management and serves as an important outcome variable in conjunction with ventricular shunt placement. However, none of the reported studies have evaluated the significance of prenatal intraventricular hemorrhage (IVH) on these important outcomes.

The incidence of IVH in ONTD has not been reported. This may be partially explained by the challenge in identification by both sonography and MR imaging. For example, sonography is less sensitive in detecting blood products and MR imaging may confuse normal germinal matrix vessels as hemorrhage, particularly on EPI sequences. Furthermore, imaging findings of IVH, including ependymal nodularity, overlap with those of gray matter heterotopia (GMH), a common finding in NTD. The aim of this study is to investigate fetal sonography and MR imaging findings of patients with ONTD with attention to prenatal IVH to determine the incidence and its associations with prenatal and postnatal HH, postnatal IVH, and ventricular shunt placement to better inform patient management and family counseling.

**MATERIALS AND METHODS**

**Setting**

As a large referral center for ONTD diagnosis and management, including fetal surgery, our institution evaluates approximately 100 new patients per year with this diagnosis. A single-institution retrospective case-control study was conducted after approval from our institutional review board. All patients who underwent evaluation for suspected NTD with fetal sonography at our institution between January 1, 2013 and April 24, 2018 were screened. Patients with a confirmed diagnosis of MMC or MS were included.

**Study Population and Case Selection**

Patients with discrepant diagnoses of the type of NTD (MMC or MS) between imaging modalities, those without fetal MR imaging, inconclusive HH, or multiple congenital anomalies or chromosomal abnormalities were excluded. The presence or absence of IVH and GMH by both prenatal sonography and MR imaging studies was used for classification. Cases of IVH were determined by the presence of IVH and the absence of GMH by both modalities (Figs 1 and 2). Controls were defined as fetuses with neither IVH nor GMH (n = 229). Studies were excluded from final analyses if there was disagreement between imaging modalities of the presence or absence of IVH or the presence of GMH to maximize the true-positive rate of IVH. Ultimately, a total of 135 subjects with findings of GMH were excluded from the final analyses.

**Data Sources**

Data were extracted from electronic medical records. If imaging reports were incomplete, variables were unavailable, or if an inconsistency in the report was discovered (ie, dictation error), images were evaluated by a pediatric radiologist for resolution and to verify data accuracy. Report data were intentionally
included in this retrospective study as it is the actionable information used to make critical treatment and management decisions during prenatal evaluation and consultation.

**Imaging Protocols**

Sonography and MR imaging examinations were performed per clinical protocols and before any surgical intervention. Detailed fetal sonography evaluation included transabdominal gray-scale and Doppler evaluation to assess the ventricular size, presence of ependymal nodularity, intraventricular debris/blood products, and the NTD. Transvaginal sonography was performed in all patients considering in utero fetal closure or for further anatomic evaluation of the brain and/or spine if transabdominal images were suboptimal. MR imaging examinations were performed on a 1.5T or 3T magnet using an 18-channel body coil in line with a spine coil with the following sequences: T2-weighted HASTE and steady-state free precession, and EPI performed in orthogonal planes through the fetal brain and spine. Postnatal MR imaging was performed with sequences including T2-weighted FSE, FLAIR, and CISS, T1-weighted FLASH or MPRAGE, gradient-echo, spin-labeled perfusion imaging, and DWI.

**Variables and Study Outcomes**

Clinical data included: fetal sex, gestational age, ventricular shunt placement within 12 months of birth, type of pregnancy (singleton or multiple), presence of an additional anomaly, and age at the time of imaging studies and surgical interventions. Eligibility for in utero closure was determined by institutional protocol and in accordance with the MOMS trial criteria, and all were performed as an open procedure. Prenatal sonography data included: gestational age (GA) by sonography, biparietal diameter (BPD), head circumference (HC), head circumference to abdominal circumference ratio (HC/AC), cerebellar diameter, transverse third ventricular size, level of the osseous defect (T12 or higher, L1–L3, or L4 or lower), transverse lateral cerebral ventricle measurement at the level of the glomus of the choroid plexus, and absolute difference in lateral ventricular size.

**Bias Assessment**

Patients were selected consecutively. All data were extracted by experienced research personnel using the Research Electronic Data Capture (REDCap) platform. Data were reviewed and validated by a board-certified radiologist with 3 or 8 years of experience. Concordance between prenatal imaging modalities identifying IVH was evaluated. Study personnel were blinded to postnatal findings of IVH at the time of data extraction and image review.

**FIG 2.** Prenatal intraventricular hemorrhage in open neural tube defect of a fetus (22 weeks and 3 days gestational age) with myelomeningocele and intraventricular hemorrhage with resolution of hindbrain herniation and intracranial blood products postnatally following in utero surgical closure. A, Sagittal gray-scale sonography image through the fetal lumbosacral spine demonstrates a myelomeningocele with a large, thin-walled sac. Transverse gray-scale sonography images through the fetal skull demonstrate (B) dilation of the third ventricle measuring 2 mm in transverse dimension and (C) layering echogenic material within the right lateral ventricle concerning for intraventricular hemorrhage. Corresponding fetal MR imaging demonstrates multiple foci of susceptibility artifact within the right lateral ventricle on (D) transverse and (E) coronal EPI through the fetal brain consistent with intraventricular hemorrhage. F, Sagittal T2WI through the fetal brain demonstrates hindbrain herniation. Following in utero surgical closure at 24 weeks and 5 days gestational age, postnatal MR imaging performed at 7 days of age demonstrates (G) resolution of hindbrain herniation on sagittal T2WI through the neonatal brain and there was no evidence of intraventricular hemorrhage on any sequence. RT, right; LT, left.
**Statistical Analysis**

R statistical software (version 3.5.1 for Windows, R Core Team) and packages ‘vcd’, ‘aod’, ‘pscl’, and ‘ggplot2’ were used. Cases and controls were compared using $\chi^2$ or Fisher exact test and independent t Student or Wilcoxon rank sum, as appropriate. Concordance was evaluated with Cohen $k$. Primary outcomes were evaluated using multivariate logistic regression models. Variables were included if univariate regression was significant (Wald test, $P < .05$), and McFadden pseudo$R^2$ was used to determine effect size. Cut-points for significant continuous predictors were established using predicted probabilities and were included in an additional model as binary predictors. Analyses were considered significant at $P < .05$.

**RESULTS**

Five hundred four cases with confirmed MMC ($n = 369, 73.2\%$) or MS ($n = 135, 26.8\%$) by prenatal sonography were included (Fig 3). Four hundred eighty-seven of 504 (96.6\%) also underwent fetal MR imaging. Overall, mean gestational age at prenatal sonography was 22 weeks and 4 days ($\pm 17$ days), and mean maternal age was 29.75 ($\pm 5.37$) years. IVH was reported prenatally in 83 of 504 (16.5\%) sonography and 108 of 487 (22.2\%) MR imaging examinations. Agreement between imaging modalities was observed in 404 of 487 cases (83\%). Of these, IVH was reported in 52 of 404 (13\%) ($\kappa = 0.454$, $P < .001$). On postnatal MR imaging, IVH was reported in 38 of 223 (17\%) neonates. Compared with postnatal MR imaging, concordance with prenatal imaging modalities was lower but remained significant ($\kappa = 0.131, P = .049$ and $\kappa = 0.223, P = .001$ for fetal sonography and MR imaging, respectively).

The On-line Table compares the findings between IVH and controls. In cases of IVH, there was a higher prevalence of HH, on both prenatal and postnatal imaging studies, though this only reached statistical significance prenatally ($P < .05$). In cases of IVH, we found significantly increased bilateral lateral ventricular size ($P < .001$), increased head biometric measurements ($P < .001$), and third ventricular diameter ($P < .001$). The level of the osseous defect was not shown to be predictive of HH in our cohort.

A multivariate model evaluating the association between prenatal IVH and HH was performed and included the presence of IVH, largest lateral ventricle by sonography, BPD, HC, GA, level of osseous defect, and transverse third ventricular diameter by sonography (Wald test, $\chi^2 = 31.8, P < .001$, pseudo$R^2 = 0.27$). IVH was more common in prenatal HH but not independently predictive (OR = 3.1 [95\% CI, 0.5–60]). Decreased BPD (OR = 0.02 [95\% CI, 0.001–0.37]), decreased HC (OR = 0.38 [95\% CI, 0.16–0.89]), increased GA (OR = 5.89 [95\% CI, 2–19]), and increased third ventricular size (OR = 8.4 [95\% CI, 2.2–37]) were predictive of prenatal HH (Table 1 and Fig 4). The level of the osseous defect was not shown to be predictive of HH in our cohort. After calculating cut-points, an additional multivariate logistic regression model showed good fit ($\chi^2 = 46.4, P < .001$).
pseudor$^2 = 0.07$) (Table 1). From this second model, only a third ventricular diameter $>1$ mm (OR $= 3.4$ [95% CI, $1.7$–$7.5$]) was independently predictive of prenatal HH. Mean third ventricular size was higher in the setting of HH $(1.15 \pm 0.74$ mm versus $0.83 \pm 0.35$ mm; $P < .001$). A multivariate model evaluating variables associated with postnatal HH was performed and included: IVH, largest ventricular size by MR imaging, GA at birth, prenatal HH, and history of in utero surgical closure (Wald test, $\chi^2 = 13.3$, $P = .02$, pseudor$^2 = 0.31$). Prenatal IVH was not predictive of postnatal HH (OR $= 2.8$ [95% CI, 0.5–16]) (Table 2). Not surprisingly, prenatal HH predicted postnatal HH (OR $11.4$ [95% CI, 2.4–75]), but a history of in utero surgical closure was negatively predictive (OR $= 0.04$ [95% CI, 0.01–0.16]).

A final multivariate regression model showed that prenatal IVH was predictive (OR $= 15.7$ [95% CI, 2.8–80]) of postnatal IVH, while a history of in utero surgical closure (OR $= 0.08$ [95% CI, 0.01–0.39]) was negatively predictive (Wald test, $\chi^2 = 25.4$, $P < .001$, pseudor$^2 = 0.24$), suggesting that fetal closure is protective against persistent IVH. No association between prenatal IVH and postnatal ventricular shunt placement was observed (On-line Table).

### DISCUSSION

IVH is commonly detected by sonography and MR imaging in prenatal evaluation of ONTD and frequently described on postnatal MR imaging. Prenatal IVH is associated with, but not independently predictive of, prenatal and postnatal HH likely related to its close association with increased ventriculomegaly. Third ventricular size was strongly associated and a 1 mm cut-point predicted prenatal HH. Postnatal HH was associated with prenatal HH and history of fetal closure was negatively predictive. These findings confirm that in utero surgical closure reverses HH, and we found that this outcome is independent of the presence of prenatal IVH. Last, prenatal IVH was not associated with a requirement for postnatal ventricular shunt placement.

Ventriculomegaly is a common finding in MMC and is thought to be secondary to obstruction of CSF by HH. In cases of open NTD, associated HH is an important factor in determining eligibility for prenatal surgical intervention and contributes to predictions of patient outcomes. We found an expected association between HH and decreased head biometric measurements as the HH limits normal calvarial expansion, which may serve as an additional clue to the presence of HH. We also found an association between HH and increased GA, suggesting that the progression of HH is possible later in gestation. Therefore, patients initially excluded from fetal surgery due to absent HH may benefit from follow-up evaluation for subsequent HH development.

In delineating the influence of prenatal IVH in our cohort, we found that ventricular size and head biometric parameters were significantly larger in cases of IVH, suggesting that hemorrhage may contribute to ventriculomegaly. Despite this association, we found that neither IVH nor increased GA, suggesting that the progression of HH is possible later in gestation. Therefore, patients initially excluded from fetal surgery due to absent HH may benefit from follow-up evaluation for subsequent HH development.

While concordance for IVH detection was moderate prenatally and reduced postnatally, overall concordance remained statistically significant. These findings may be explained by the self-limited nature of IVH, time-associated factors, the potential for interval hemorrhage, and unavoidable differences in the timing of imaging. Our results are similar to those reported comparing prenatal and postnatal imaging studies for GMH. In addition, our multivariate model shows that fetal closure may even promote IVH resolution; we suspect by reversal of HH and improvement in CSF flow dynamics thus allowing clearance of blood products. This is concordant with the finding that fetal closure is protective against persistent IVH.
with results reporting a higher prevalence of IVH in infants who did not undergo in utero surgery. Ultimately previously considered possible exclusion criteria, prenatal IVH should not preclude eligibility for in utero closure, and families should be counseled and treated accordingly.

The limitations of this study are the retrospective nature and decision to use imaging report data, though this accounts for variability in clinical practice. We ran concordance analyses among imaging studies before case classification and defined IVH based on findings by both sonography and MR imaging to account for this bias and increase data reliability. Previous studies have implemented a grading system in the severity of HH, which would be of benefit for more detailed characterization, but the severity has not been shown to significantly change without fetal closure, does not influence patient counseling or management at our institution, and would be challenging to implement given the variability in the timing of postnatal MR imaging in our cohort; but could be a source of further study. Lastly, subtle changes in sonography and MR image acquisition over the study period may affect the study results, though performing statistical analyses as case-control series may decrease any potential influence on outcomes.

CONCLUSIONS

IVH is prevalent in the prenatal evaluation of fetuses with ONTD. While IVH is associated with prenatal HH, this is not independent and seen in conjunction with increased ventricular size. The transverse third ventricular diameter appears to be a good radiologic marker of prenatal HH and, if larger than 1 mm, it is strongly predictive. The presence of prenatal IVH does not prevent resolution of HH following fetal surgical closure and does not increase the need for postnatal shunt placement. Overall, our results support that prenatal IVH in ONTD is self-limited, particularly in those with fetal closure that is protective against postnatal IVH, likely secondary to the restoration of CSF flow dynamics and clearance of blood products.

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