Congenital abnormalities of the posterior fossa

Bosemani, Thangamadhan; Orman, Gunes; Boltshauser, Eugen; Tekes, Aylin; Huisman, Thierry A G M; Poretti, Andrea

Abstract: The frequency and importance of the evaluation of the posterior fossa have increased significantly over the past 20 years owing to advances in neuroimaging. Nowadays, conventional and advanced neuroimaging techniques allow detailed evaluation of the complex anatomic structures within the posterior fossa. A wide spectrum of congenital abnormalities has been demonstrated, including malformations (anomalies due to an alteration of the primary developmental program caused by a genetic defect) and disruptions (anomalies due to the breakdown of a structure that had a normal developmental potential). Familiarity with the spectrum of congenital posterior fossa anomalies and their well-defined diagnostic criteria is crucial for optimal therapy, an accurate prognosis, and correct genetic counseling. The authors discuss the spectrum of posterior fossa malformations and disruptions, with emphasis on neuroimaging findings (including diagnostic criteria), neurologic presentation, systemic involvement, prognosis, and risk of recurrence.

DOI: [https://doi.org/10.1148/rg.351140038](https://doi.org/10.1148/rg.351140038)

Posted at the Zurich Open Repository and Archive, University of Zurich

ZORA URL: [https://doi.org/10.5167/uzh-112443](https://doi.org/10.5167/uzh-112443)

Published Version

Originally published at:


DOI: [https://doi.org/10.1148/rg.351140038](https://doi.org/10.1148/rg.351140038)
Congenital Abnormalities of the Posterior Fossa

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Introduction

Over the past 2 decades, significant advances in pre- and postnatal neuroimaging techniques, the development of next-generation genetic sequencing, and an increase in animal model research have led to more accurate definition and classification of congenital abnormalities of the posterior fossa and a better understanding of their pathogenesis. Classifications of congenital posterior fossa abnormalities based on neuroimaging, molecular genetic, and developmental biologic criteria have been proposed (1–3).

Congenital posterior fossa anomalies may result from inherited (genetic) or acquired (disruptive) causes (4,5). A malformation is defined as a congenital morphologic anomaly of a single organ or body part due to an alteration of the primary developmental program caused by a genetic defect (6). Gene mutations causing malformations may be de novo (ie, new in the affected child, rather than present in or transmitted by the parents) or inherited from the parents. Inherited mutations are transmitted in different patterns (eg, autosomal recessive, X linked) with differences in recurrence risk for further offspring and for potential transmission of mutations from the patients to their children. The latter is the case for autosomal recessive–inherited diseases (from an unaffected mother or father to her or his child), autosomal dominant–inherited diseases (from an affected mother or father to her or his child), X-linked diseases (from an affected mother to her son), and mitochondrially inherited diseases (from an affected mother to her child). A disruption is defined as a congenital morphologic anomaly caused by the breakdown of an anatomic structure that had a normal developmental potential (6). Disruptive causes include prenatal infection, hemorrhage, and ischemia, among others. Disruptions are acquired lesions without risk of...
TEACHING POINTS

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- Each part of the cerebellum (vermis and hemispheres) may be hypoplastic and/or dysplastic, resulting in global cerebellar involvement or predominantly vermian involvement. Predominant involvement of the cerebellar hemispheres is uncommon and is characteristic of pontocerebellar hypoplasia as defined by Barth, as well as disruptive cerebellar development in very premature newborns. Typically, posterior fossa malformations involve both cerebellar hemispheres equally. Hypoplasia and/or dysplasia of only one cerebellar hemisphere is most likely a sequel of a prenatal disruptive lesion such as hemorrhage.

- To classify malformations that do not meet the criteria for DWM, terms such as “Dandy-Walker variant,” “Dandy-Walker complex,” or “Dandy-Walker spectrum” have been introduced in the past. However, these terms lack specificity and can create considerable confusion. Accordingly, these terms should not be used, and a more detailed anatomic descriptor (eg, interior cerebellar vermis or global cerebellar hypoplasia) is to be preferred.

- The “molar tooth sign” is the diagnostic criterion for Joubert syndrome and consists of elongated, thickened, and horizontally oriented superior cerebellar peduncles; a deep interpontine fossa; and vermian hypoplasia.

- Cerebellar injury occurs in up to 20% of preterm infants prior to 32 weeks gestation. Impaired cerebellar development in preterm newborns may be caused by primarily destructive injuries or secondary lesions. Primarily destructive injuries include hemorrhage and ischemia, and cause cerebellar tissue loss. Secondary lesions lead to underdevelopment of the cerebellum and are caused by direct (eg, hemosiderin–blood products, glucocorticoid exposure, undernutrition) or indirect (impaired trans-synaptic trophic effects) effects on the cerebellum.

- A disruption is defined as a congenital morphologic anomaly caused by the breakdown of an anatomic structure that had a normal developmental potential. Disruptive causes include prenatal infection, hemorrhage, and ischemia, among others. Disruptions are acquired lesions without risk of recurrence. However, a genetic predisposition to disruptive lesions such as dominant mutations in COL4A1 may be present.

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- Accurate diagnoses of these complex abnormalities are of paramount significance for three primary reasons: to determine (a) inheritance pattern and risk of recurrence, (b) involvement of other systems (eg, kidneys and liver), and (c) prognostic implications for the child and his or her family. There are major implications for genetic counseling and family planning secondary to the significant difference in recurrence risk between a malformation and a disruption. Neuroimaging plays a key role in the diagnosis of posterior fossa abnormalities, and the challenge for the neuroradiologist is to provide the clinician with precise and up-to-date diagnostic information.

In this article, we discuss congenital posterior fossa abnormalities in terms of the normal anatomy of the posterior fossa, imaging techniques and strategy, and the most important congenital abnormalities of the posterior fossa, making use of a simple classification scheme based on the neuroimaging-based scheme proposed by Doherty et al (8). For each disorder, we summarize the key neuroimaging findings that are needed for the diagnosis and review the pattern of inheritance, systemic involvement, and long-term neurologic prognosis to emphasize the high clinical relevance of an accurate diagnosis. Although cerebellar agenesis and global cerebellar hypoplasia may result from either a malformation or a disruption, we discuss these disease entities only under “Cerebellar Disruptions” to avoid overlap.

**Normal Posterior Fossa Anatomy**

Conventional magnetic resonance (MR) imaging allows detailed evaluation of the anatomy of the posterior fossa and its contents (8). A midline sagittal T1- or T2-weighted sequence is ideal for showing the size of the posterior fossa, the shape and size of the vermis, and the size and morphology of the fourth ventricle and brainstem (Fig 1a). The vermis is divided into three parts by the primary and prepyramidal fissures. The rostrocaudal length of the ventral pons should be approximately twice that of the midbrain from the isthmus (ventral midbrain–pons junction) to the third ventricle, whereas the rostrocaudal length of the midbrain should be roughly the same as that of the medulla (from the level of the obex to the level of the ventral pontomedullary junction) (8). The posterior margin of the brainstem extending from the caudal aqueduct to the obex should be a straight line. The fastigium, or summit of the fourth ventricle, should lie just below the midpoint of the ventral pons on sagittal images. The cerebellar hemispheres and peduncles can be assessed on parasagittal images, whereas the size and morphology of the vermis, cerebellar hemispheres, dentate nuclei, and superior and middle cerebellar peduncles can be evaluated on axial images. In addition, the cerebellar folia run parallel to the calvaria (onion-like configuration) (Fig 1b). The individual cerebellar folia are grouped together as lobules arising from a common ray of medullary white matter. The 10 lobules of the
Figure 1. Normal anatomy of the posterior fossa in a 17-year-old girl. (a) Midsagittal T1-weighted MR image shows a normal-sized posterior fossa, a normal vermis, an appropriate-sized pons (rostrocaudal length approximately twice that of the midbrain and medulla), a flat dorsal surface of the brainstem (line), a normal position of the fastigium just below the midpoint of the ventral pons (black arrowhead), and primary (arrow) and prepyramidal (white arrowhead) fissures separating the vermis into three segments. (b) Axial T2-weighted MR image of the posterior fossa shows normal orientation of the cerebellar folia, which run parallel to the calvaria ("onion-like" orientation) (arrows). (c) Coronal T2-weighted MR image of the posterior fossa shows cerebellar fissures radiating toward the nuclei (arrows). (d) Axial color-coded fractional anisotropic map obtained at the level of the lower pons shows the middle cerebellar peduncles (anteroposterior orientation) (green) lateral to the brainstem (white arrows), anteriorly situated corticospinal tracts (superoinferior orientation) (blue) (black arrows), and posteriorly situated medial lemnisci (arrowheads), with the transverse pontine fibers (transverse orientation) in between (red (+)). (e) Axial color-coded fractional anisotropic map obtained at the level of the pontomesencephalic junction shows the superior cerebellar peduncles (green-blue) (arrows) and transverse fibers at the decussation of the ventral tegmentum and superior cerebellar peduncles (red) (arrowhead).

cerebellar hemispheres are divided into three major lobes: the anterior lobe (lobules I–V), the posterior lobe (lobules VI–IX), and the flocculonodular lobe (lobule X). The anterior lobe is separated from the posterior lobe by the primary fissure, and the posterior lobe is separated from the flocculonodular lobe by the posterolateral fissure. Coronal images show fissures radiating toward the cerebellar nuclei (Fig 1c). In addition, the superior, middle, and inferior cerebellar peduncles can be assessed in terms of size, symmetry, contour, and location.
Imaging Techniques and Strategy
Ultrasonography (US), computed tomography, and MR imaging (the only modality discussed in this article) are the key neuroimaging modalities for the evaluation of the neonatal and pediatric posterior fossa. Isotropic three-dimensional T1-weighted images allow good anatomic evaluation in three imaging planes in a reasonable amount of time. High-resolution T2-weighted images are ideal for more detailed evaluation of cerebellar folia, interfolial spaces, cerebellar cortex, gray matter–white matter differentiation, dentate nuclei, and brainstem cranial nerve nuclei. Depending on the clinical presentation, suspected diagnosis, or pathologic condition, the use of various additional sequences may be considered. Diffusion tensor imaging provides detailed qualitative and quantitative information about the microstructure and organization of the white matter tracts within the posterior fossa (Fig 1d, 1e) (9). Susceptibility-weighted imaging is highly sensitive for blood, blood products, and calcifications and may be helpful in supporting the notion of a disruptive pathomechanism (10). The indications for the essential anatomic and functional MR imaging sequences to be included in a detailed neuroimaging study of congenital posterior fossa anomalies are summarized in Table 1.

<table>
<thead>
<tr>
<th>Sequence</th>
<th>Plane</th>
<th>Key Applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>3D T1W</td>
<td>Axial, coronal, sagittal</td>
<td>Excellent differentiation between gray matter and white matter, acquisition of high-resolution anatomic information</td>
</tr>
<tr>
<td>T2W</td>
<td>Axial, coronal</td>
<td>Acquisition of high-resolution anatomic information; delineation of cortex, white matter, and gray matter nuclei</td>
</tr>
<tr>
<td>Diffusion tensor</td>
<td>Axial</td>
<td>Evaluation of white matter microstructural integrity, identification of white matter tracts</td>
</tr>
<tr>
<td>CISS*</td>
<td>Axial + MPR</td>
<td>Evaluation of cerebellar folia, cranial nerves, ventricles, and foramina</td>
</tr>
<tr>
<td>Susceptibility-weighted</td>
<td>Axial</td>
<td>Identification and characterization of hemorrhage, blood products, calcification, and iron accumulation</td>
</tr>
</tbody>
</table>

Note.—CISS = constructive interference in the steady state, MPR = multiplanar reformation, 3D = three-dimensional, T1W = T1-weighted, T2W = T2-weighted.
*Optional (depending on indication).

Malformations
As mentioned earlier, we classify posterior fossa malformations using a simple classification scheme based on the neuroimaging pattern proposed by Doherty et al (8). We divide these malformations into (a) predominantly cerebellar, (b) cerebellar and brainstem, (c) predominantly brainstem, and (d) predominantly midbrain malformations (Table 2). Although such a classification scheme may not be as comprehensive as a genetics- or embryology-based classification scheme, we are convinced that it is practical and accessible for daily clinical practice.

Predominantly Cerebellar Malformations
A malformed cerebellum may be hypoplastic (reduced cerebellar volume), dysplastic (abnormal cerebellar foliation, fissuration, and architecture of the cerebellar white matter), or hypodysplastic (combination of hypoplasia and dysplasia). Each part of the cerebellum (vermis and hemispheres) may be hypoplastic and/or dysplastic, resulting in global cerebellar involvement or predominantly vermian involvement. Predominant involvement of the cerebellar hemispheres is uncommon and is characteristic of pontocerebellar hypoplasia (PCH) as defined by Barth (11), as well as disruptive cerebellar development in very premature newborns. Typically, posterior fossa malformations involve both cerebellar hemispheres equally. Hypoplasia and/or dysplasia of only one cerebellar hemisphere is most likely a sequela of a prenatal disruptive lesion such as hemorrhage. In this section, besides DWM, we discuss other cystic posterior fossa malformations (Blake pouch cyst,
Dandy-Walker Malformation.—DWM is the most common posterior fossa malformation and typically occurs only sporadically, with a low overall risk of recurrence (1%–5%) (8). DWM may be isolated or may occur as part of chromosomal anomalies or well-defined Mendelian disorders such as Ritscher-Schinzel or cranio-cerebello-cardiac syndrome (Online Mendelian Inheritance in Man [OMIM] 220210). In addition, rare mutations in six genes (ZIC1, ZIC4, FOXC1, FGF17, LAMC1, and NID1) have been found in a few patients with DWM. On the basis of the function of these genes, DWM may represent a complex disruption of the interaction between the developing cerebellum and the developing posterior fossa mesenchyme and its derivates.

The majority of patients with DWM present before 1 year of age with signs and symptoms of increased intracranial pressure (12). Macrocephaly is the most common manifestation, affecting 90%–100% of children during the first months of life (13). Nowadays, the diagnosis of DWM is made prenatally in an increasing number of patients. The outcome of children with DWM is variable. Limited evidence shows that normal lobulation of the cerebellar vermis and the absence of associated brain abnormalities (particularly callosal dysgenesis) may be prognostic for normal cognitive function (14,15). Systemic involvement (eg, cardiovascular, urogenital, or skeletal anomalies) may be seen in DWM, is associated with defined syndromes, and also influences the prognosis (poorer outcome).

The key neuroimaging features of DWM are (a) hypoplasia (or, rarely, agenesis) of the cerebellar vermis (whose inferior portion is typically affected, possibly in combination with its superior portion), which is elevated and upwardly rotated; and (b) dilatation of the cystic-appearing fourth ventricle, which consequently may fill the entire posterior fossa (Fig 2a). Both findings are consistently present and are required for the diagnosis. The cystic structure within the posterior fossa is not the posterior fossa subarachnoid space or an arachnoid cyst; instead, it represents the cystlike dilatation of the fourth ventricle secondary to hypoplasia of the inferior cerebellar vermis. The cerebellar hemispheres are typically displaced anterolaterally, but their size and morphology are usually normal (9). The posterior fossa is usually enlarged, and the tentorium as well as the torcular and transverse sinuses are elevated (2).

### Table 2: Neuroimaging-based Differential Diagnosis for Congenital Posterior Fossa Abnormalities

<table>
<thead>
<tr>
<th>Neuroimaging Pattern</th>
<th>Malformations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Predominantly cerebellar involvement</td>
<td>Predominantly vermian, DWM and other cystic lesions of the posterior fossa, including Blake pouch cyst, mega cisterna magna, and arachnoid cysts; isolated vermic hypoplasia; rhombencephalosynapsis</td>
</tr>
<tr>
<td>Predominantly vermian</td>
<td>DWM and other cystic lesions of the posterior fossa, including Blake pouch cyst, mega cisterna magna, and arachnoid cysts; isolated vermic hypoplasia; rhombencephalosynapsis</td>
</tr>
<tr>
<td>Global cerebellar</td>
<td>Malformations of cortical development: lissencephaly (RELN, VLDR1, tubulin genes), polymicrogyria (tubulin genes, GPR56), periventricular nodular heterotopia (FLNA), and primary microcephaly; macrocerebellum; cerebellar dysplasia, including Chudley-McCullough syndrome; global cerebellar hypoplasia in multiple genetic syndromes (eg, Ritscher-Schinzel syndrome, CHARGE syndrome, Delleman syndrome); congenital CMV infection</td>
</tr>
<tr>
<td>Unilateral cerebellar</td>
<td>Isolated unilateral cerebellar hypoplasia, PHACES syndrome, COL4A1 mutations, cerebellar cleft</td>
</tr>
<tr>
<td>Cerebellar and brainstem involvement</td>
<td>PCH type 1–10; glycosylation type 1a congenital disorder; CASK mutations; Joubert syndrome; cerebellar agenesis caused by a disruptive lesion or representing a malformation due to PTF1A mutations; congenital muscular dystrophies due to defective dystroglycan O-glycosylation; vanishing cerebellum in Chiari 2 malformation; cerebellar disruption secondary to prematurity</td>
</tr>
<tr>
<td>Predominantly brainstem involvement</td>
<td>Pontine tegmental cap dysplasia; horizontal gaze palsy and progressive scoliosis (ROBO3 mutations)</td>
</tr>
<tr>
<td>Predominantly midbrain involvement</td>
<td>Dysplasia of the diencephalic-mesencephalic junction</td>
</tr>
</tbody>
</table>

Note.—CHARGE = coloboma, heart defect, choanal atresia, retarded growth and development, genital abnormality, and ear abnormality, CMV = cytomegalovirus, DWM = Dandy-Walker malformation, PCH = pontocerebellar hypoplasia.
Figure 2. Cystic malformations of the posterior fossa as seen at sagittal T1-weighted MR imaging. (a) Image shows a DWM with a hypoplastic vermis in upward rotation (arrow), cystic dilatation of the fourth ventricle (white arrow) with posterior extension and communication with an enlarged posterior fossa, and supratentorial hydrocephalus (arrowhead). (b) Image shows a Blake pouch cyst with enlargement of the fourth ventricle, which communicates with an infravermian cystic compartment (arrow) corresponding to enlargement of the Blake pouch; a normal vermis; and supratentorial hydrocephalus (arrowhead). (Reprinted, with permission, from reference 19.) (c) Image shows mega cisterna magna (arrow), a normal vermis, a normal fourth ventricle, an enlarged posterior fossa, scalloping of the occipital bone, and the absence of hydrocephalus. (d) Image shows a retrocerebellar arachnoid cyst that is isointense relative to CSF (white arrows), with apparent enlargement of the posterior fossa, scalloping of the occipital bone, mass effect on the dorsal aspect of a normal-appearing vermis (black arrow), a normal fourth ventricle, and supratentorial hydrocephalus (arrowhead). (e) Image shows isolated inferior vermian hypoplasia with partial absence of the inferior vermis (arrow) and apparent but not true enlargement of the fourth ventricle.

In fact, the displacement of the cerebellar hemispheres and the upward displacement of the tentorium, torcular herophili, and transverse dural sinuses are caused by the cystlike dilatation of the fourth ventricle and result in global enlargement of the posterior fossa. Additional malformations, including dysgenesis or agenesis of the corpus callosum, occipital encephalocele, polymicrogyria, and heterotopia, may be present in 30%–50% of individuals (2). Hydrocephalus is associated with DWM in about 90% of patients (2). Cerebrospinal fluid (CSF) flow studies may be added to better understand the complex hydrodynamic changes in patients with DWM. If aqueductal stenosis or occlusion is shown with CSF flow...
studies, cystoperitoneal shunt placement is not sufficient and placement of a ventriculoperitoneal shunt is required.

Neuroimaging diagnostic criteria usually make it possible to distinguish DWM from other cystic posterior fossa malformations (Table 3). In some patients, however, only a partial neuroimaging phenotype of DWM is noted. To classify malformations that do not meet the criteria for DWM, terms such as “Dandy-Walker variant,” “Dandy-Walker complex,” or “Dandy-Walker spectrum” have been introduced in the past. However, these terms lack specificity and can create considerable confusion. Accordingly, these terms should not be used, and a more detailed anatomic descriptor (eg, inferior cerebellar vermis or global cerebellar hypoplasia) is to be preferred (2).

**Blake Pouch Cyst.**—Blake pouch cyst is caused by lack of fenestration of the Blake pouch, resulting in absence of communication between the fourth ventricle and the subarachnoid space (16–18) and leading to tetraventricular hydrocephalus. The cerebellum has a normal size and shape. Blake pouch cyst occurs sporadically, and no recurrence risk has been reported.

Hydrocephalus and macrocephaly are the most common presenting features in the neonatal period (18). The long-term outcome depends mostly on complications related to neurosurgical procedures. In the absence of shunt-related complications, the prognosis is generally favorable.

Typical neuroimaging findings include the presence of a cyst in a retrocerebellar or infratentorial location, which is essentially a diverticulum of the consequently enlarged fourth ventricle (Fig 2b). The presence of this cyst is responsible for the displacement of the choroid plexus inferior to the vermis along the anterosuperior aspect of the cyst. The displacement of the choroid plexus is best visualized as an enhancing structure on sagittal contrast material–enhanced T1-weighted images (17). The consistent presence of hydrocephalus allows the differentiation of Blake pouch cyst from mega cisterna magna. Mild mass effect may result in indentation of the inferior vermis or of the caudal and medial aspects of the cerebellar hemispheres. The posterior fossa is typically normal in size. Supratentorial morphologic abnormalities other than hydrocephalus are usually absent.

**Mega Cisterna Magna.**—Mega cisterna magna is an enlarged cisterna magna (≥10 mm on mid-sagittal images) with an intact vermis, a normal fourth ventricle, and, in some patients, an enlarged posterior fossa (Fig 2c) (16,20,21). It represents a truly focal enlargement of the subarachnoid space in the inferior and posterior portions of the posterior fossa. Consequently, mega cisterna magna freely communicates with the fourth ventricle and the cervical subarachnoid space, which is confirmed with CSF flow studies (20) and results in consistent absence of hydrocephalus. It has been suggested that delayed fenestration of the Blake pouch results in mega cisterna magna, whereas the absence of fenestration of the Blake pouch results in Blake pouch cyst (17). There is no reported recurrence risk.

Mega cisterna magna is typically an incidental finding and represents a normal variant. Accordingly, no follow-up neuroimaging studies are required. The vast majority (90%–95%) of children with isolated mega cisterna magna develop normally (15).

Neuroimaging plays a key role in identifying mega cisterna magna and in differentiating it from isolated inferior vermian hypoplasia and Blake pouch cyst. In mega cisterna magna, the presence of a normal vermis and the absence of

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### Table 3: Key Neuroimaging Features of DWM versus Other Cystic Posterior Fossa Malformations

<table>
<thead>
<tr>
<th>Malformation</th>
<th>Vermis</th>
<th>Fourth Ventricle</th>
<th>Posterior Fossa</th>
<th>Hydrocephalus</th>
<th>Occipital Bone Scalloping*</th>
</tr>
</thead>
<tbody>
<tr>
<td>DWM</td>
<td>Hypoplastic</td>
<td>Enlarged</td>
<td>Enlarged</td>
<td>Yes (in most patients)</td>
<td>No</td>
</tr>
<tr>
<td>IVH (inferior portion)</td>
<td>Hypoplastic</td>
<td>Enlarged</td>
<td>Normal</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>BPC</td>
<td>Normal</td>
<td>Enlarged</td>
<td>Normal</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>MCM</td>
<td>Normal</td>
<td>Normal</td>
<td>Inconsistently enlarged</td>
<td>No</td>
<td>Possible</td>
</tr>
<tr>
<td>PFAC</td>
<td>Normal</td>
<td>Normal or reduced</td>
<td>Normal</td>
<td>Possible</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Note.—BPC = Blake pouch cyst, IVH = isolated vermian hypoplasia, MCM = mega cisterna magna, PFAC = posterior fossa arachnoid cyst.

*Not typically seen in neonates and infants.
hydrocephalus help differentiate it from isolated inferior vermian hypoplasia and Blake pouch cyst, respectively.

**Posterior Fossa Arachnoid Cysts.**—Duplications of the arachnoid membrane produce fluid-filled cysts known as arachnoid cysts. About 10% of arachnoid cysts in children occur in the posterior fossa and may be located inferior or posterior to the vermis in a midsagittal location (retrocerebellar), cranial to the vermis in the tentorial hiatus (supravermian), anterior or lateral to the cerebellar hemispheres, or anterior to the brainstem (22). Arachnoid cysts do not communicate with the fourth ventricle or the subarachnoid space. There is no reported recurrence risk.

Macrocephaly, signs of increased intracranial pressure, and developmental delay are possible clinical findings in children with posterior fossa arachnoid cysts, particularly if CSF flow is obstructed (23). However, these cysts may be asymptomatic and discovered incidentally. Overall, children who undergo surgery and shunt placement have a favorable outcome (4,23).

Neuroimaging reveals a well-circumscribed extraxial fluid collection or cyst that is isointense relative to CSF with all sequences (Fig 2d). The presence of proteinaceous content may lead to lack of complete signal suppression with a fluid-attenuated inversion-recovery sequence. Diffusion-weighted imaging reveals free water motion or facilitated diffusion similar to that seen in CSF. The cyst walls are usually too thin to be visualized at MR imaging. Arachnoid cysts may enlarge during infancy and produce mass effect on the cerebellum and vermis, which may cause a secondary obstruction of the ventricular system, hydrocephalus, and/or remodeling or thinning of the overlying occipital bone.

**Isolated Inferior Vermian Hypoplasia.**—Isolated inferior vermian hypoplasia is a pathologic condition that is characterized by partial absence of the inferior portion of the cerebellar vermis. In the literature, there is some confusion about isolated inferior vermian hypoplasia, with some authors referring to it as Dandy-Walker variant (24). As discussed earlier, the term Dandy-Walker variant lacks specificity, creates confusion, and can lead to misdiagnosis and incorrect genetic counseling. Inferior vermian hypoplasia in isolation has no recurrence risk; however, this condition may occur in the context of Mendelian syndromes, in which case it does carry a recurrence risk.

More than 75% of patients with isolated inferior vermian hypoplasia have a favorable outcome (25,26). In some patients, mild functional deficits in fine motor activity and receptive language may be present.

The key neuroimaging finding in isolated inferior vermian hypoplasia is partial absence of the inferior vermis, a finding that is best seen on the midsagittal image (Fig 2e). The remainder of the vermis, as well as the cerebellar hemispheres, the fourth ventricle, and the posterior fossa, have a normal size and architecture. The prenatal diagnosis of isolated inferior vermian hypoplasia is reliable after 18–20 weeks gestation (25). Before 18 weeks gestation, incomplete caudal growth of the inferior vermis over the fourth ventricle may be physiologic. The prenatal US diagnosis of inferior vermian hypoplasia is not reliable. Fetal MR imaging provides a more sensitive and specific diagnosis of inferior vermian hypoplasia compared with US, but the false-positive rate remains high (~30%) (25).

**Cerebellar Involvement in Migrational Disorders.**—Posterior fossa abnormalities including hypoplasia of the pons and cerebellum have been reported in migrational disorders (27). A severely hypoplastic cerebellum with a markedly reduced foliation pattern and severe hypoplasia of the pons are characteristic findings in patients with mutations in RELN (28) and VLDLR (29). Typically, the cerebellar vermis is more affected than the hemispheres. Patients with autosomal recessive mutations in RELN typically present with microcephaly, seizures, marked cognitive impairment, and congenital lymphedema. The supratentorial brain shows a simplified gyration pattern in the majority of patients. Patients with autosomal recessive mutations in VLDLR have a similar but milder posterior fossa neuroimaging phenotype compared with patients with mutations in RELN.

Mutations in a number of neuron-specific α- and β-tubulin genes (eg, TUBA1A, TUBB2B) are associated with a spectrum of posterior fossa abnormalities ranging from severe PCH to mild cerebellar hypoplasia, pontine cleft, and/or asymmetric cerebral peduncles (Fig 3a) (30). In addition, asymmetry of the cerebellar hemispheres and peduncles may be present. Dysmorphic basal ganglia with an abnormal internal capsule are a consistent and characteristic supratentorial finding (Fig 3b). Migrational abnormalities ranging from lissencephaly to polymicrogyria and callosal hypoplasia may also be present. The majority of mutations in tubulin genes are de novo.

**Rhombencephalosynapsis.**—Rhombencephalosynapsis is characterized by absence of the vermis and continuity of the cerebellar hemispheres, dentate nuclei, and superior cerebellar peduncles (31). The majority of patients are nonsyndromic. However, rhombencephalosynapsis is a key feature of Gómez-López-Hernández syndrome.
(OMIM 601853 [parietal alopecia, trigeminal anesthesia, and craniofacial dysmorphic signs]) (32,33) and may be seen in patients with associated VACTERL (vertebral anomalies, anal atresia, cardiovascular anomalies, tracheoesophageal fistula, renal anomalies, and limb defects) (34). The sporadic nature of rhombencephalosynapsis contributes to its low recurrence risk.

The most common clinical manifestations of rhombencephalosynapsis include truncal and/or limb ataxia, abnormal eye movements, head stereotypies, and delayed motor development. Long-term cognitive outcome varies from severe impairment to normalcy (32). Patients with more severe rhombencephalosynapsis, associated holoprosencephaly, or VACTERL features have more severe neurodevelopmental impairment (34).

Systemic involvement is uncommon. The key neuroimaging findings in rhombencephalosynapsis are agenesis or hypogenesis of the vermis and continuity (often called fusion) of the cerebellar hemispheres, superior cerebellar peduncles, and dentate nuclei, which creates a horseshoe-shaped arch across the midline, resulting in a keyhole-shaped fourth ventricle (Fig 4). Posterior coronal T2-weighted images are crucial in showing the horizontal folial pattern (Fig 4a), whereas the midsagittal T1-weighted image depicts the dentate nuclei. (In normal anatomy, the vermis separates the dentate nuclei in the midline.) Rhombencephalosynapsis may be associated with other central nervous system anomalies such as hydrocephalus, mostly due to aqueductal stenosis and forebrain abnormalities including absent olfactory bulbs, dysgenesis of the corpus callosum, and absent septum pellucidum (34).

**Macrocerebellum.**—Macrocerebellum, or cerebellar hyperplasia, is an extremely rare condition that is characterized by an abnormally large cerebellum with preservation of its overall shape (35). Macrocerebellum may be isolated (non-syndromic) or part of well-defined syndromes (eg, Costello syndrome, Sotos syndrome) or neurometabolic diseases (eg, fucosidosis, mucopolysaccharidosis type I and II) (35,36). The significant difference in pathogenesis between these disorders suggests that macrocerebellum may not be a nosologic entity, but instead represents the structural manifestation of a deeper, more basic biologic disturbance common to heterogeneous diseases. The recurrence risk of macrocerebellum depends on the underlying disorder. In patients with isolated macrocerebellum, the recurrence risk is low (35).

The clinical presentation of patients with macrocerebellum is highly heterogeneous and depends mostly on the underlying disease. Ataxia, hypotonia, intellectual disability, and ocular movement disorders are common in patients with isolated macrocerebellum.

The key neuroimaging finding of macrocerebellum is disproportionate cerebellar enlargement with preserved architecture and shape, which is best seen on sagittal or axial images (Fig 5) (35,37). The cerebellar hemispheres are more affected than the vermis and may expand into adjacent anatomic regions by wrapping around the...
brainstem or herniating upward or downward. Supratentorial findings such as ventriculomegaly or white matter signal abnormalities may be present depending on the underlying disease.

**Cerebellar Dysplasia.**—*Cerebellar dysplasia* is a descriptive anatomic term. This pathologic condition includes a heterogeneous distribution of different causes, both malformations and disruptions, and hence the variable clinical presentations, additional neuroimaging findings, and recurrence risks.

The clinical manifestation of cerebellar dysplasia is highly variable, ranging from an incidental finding to severe neurologic impairment.

The key neuroimaging findings of cerebellar dysplasia are abnormal foliation, fissuration (including defective, enlarged, or vertical fissures), white matter arborization, and an obscured gray matter–white matter junction (Fig 6) (38). Findings
may be focal or global. Focal findings suggest a disruptive lesion, whereas global findings suggest a malformation. Supratentorial findings, including migrational abnormalities, callosal dysgenesis, and clastic lesions, may also be present (38,39).

Cerebellar dysplasia in the inferior cerebellar hemispheres is a consistent finding in Chudley-McCullough syndrome (OMIM 604213), an autosomal recessive disorder (8). Additional features of Chudley-McCullough syndrome include severe sensorineural hearing loss, mild developmental delay, partial agenesis of the corpus callosum, and periventricular heterotopia (40). Mutations in GPSM2 have recently been identified as the cause of Chudley-McCullough syndrome (41). This gene encodes a GTPase regulator needed for correct orientation of stem cell division (41).

**Cerebellar and Brainstem Malformations**

**Pontocerebellar Hypoplasia.**—As conceptualized by Barth, PCH is a group of autosomal recessive neurodegenerative disorders with a prenatal onset and is characterized by hypoplasia of the cerebellum and pons (11,42). To date, 10 subtypes of PCH with different phenotypes and pathogeneses (types 3 and 8 have a nonprogressive course) have been identified (Table 4) (43–46).

Coronal T2-weighted and sagittal T1- or T2-weighted imaging is essential in the evaluation of PCH (Fig 7). The cerebellar abnormality is characterized by hypoplasia with superimposed atrophy with prenatal onset. In some cases of PCH (particularly type 2, which is caused by mutations in TSEN54), there is more severe involvement of the cerebellar hemispheres. On coronal T2-weighted images, this involvement has a “dragonfly” appearance that is created by flattened cerebellar hemispheres (the “wings”) and a relatively preserved vermis (the “body”) (Fig 7b) (43).

<table>
<thead>
<tr>
<th>Subtype</th>
<th>Main Features</th>
<th>Genotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Involvement of the anterior horn cells</td>
<td>VRK1, TSEN54, RARS2, EXOSC3</td>
</tr>
<tr>
<td>2</td>
<td>Chorea/dystonia; rarely, spasticity only</td>
<td>TSEN54 (vast majority of cases); TSEN2, TSEN34</td>
</tr>
<tr>
<td>3</td>
<td>Optic atrophy, nonprogressive course</td>
<td>Locus on 7q11–21</td>
</tr>
<tr>
<td>4</td>
<td>Severe neonatal course with hypertonia and hypoventilation, absent inferior olivary prominence at autopsy</td>
<td>TSEN54</td>
</tr>
<tr>
<td>5</td>
<td>Lethal course with neonatal death, more pronounced involvement of the vermis than of the hemispheres</td>
<td>TSEN54</td>
</tr>
<tr>
<td>6</td>
<td>Lactic acidemia, high lactate level in CSF</td>
<td>RARS2</td>
</tr>
<tr>
<td>7</td>
<td>Micropenis with no palpable gonads</td>
<td>Unknown</td>
</tr>
<tr>
<td>8</td>
<td>Proportionate involvement of the vermis and hemispheres, nonprogressive course</td>
<td>CHMP1A</td>
</tr>
<tr>
<td>9</td>
<td>Dysgenesis of the corpus callosum</td>
<td>AMPD2</td>
</tr>
<tr>
<td>10</td>
<td>Dymorphic features and axonal sensorimotor neuropathy (inconsistent)</td>
<td>CLP1</td>
</tr>
</tbody>
</table>

**Table 4: Subtypes of PCH**

Figure 6. Cerebellar dysplasia in a 17-year-old girl who presented with intellectual disability and seizures. Axial T2-weighted MR image shows abnormal cerebellar foliation and fissuration with loss of the normal white matter architecture in the inferior aspect of the cerebellar hemispheres (arrows).
Figure 7. PCH type 2 in a 1-year-old child who presented with postnatal progressive microcephaly and dyskinetic movement disorders. (a) Sagittal T1-weighted MR image shows hypoplasia of the pons (arrow) and cerebellar vermis. (b) Coronal T2-weighted MR image shows reduction in the size of the cerebellar hemispheres (arrows) with relative preservation of the midline vermis, resulting in a characteristic dragonfly appearance.

Absence or significant reduction of the pontine prominence is characteristic of a mid-hindbrain disease with prenatal onset, including malformations, disruptions, or degenerative disorders (47). It is remarkable that hereditary progressive cerebellar disorders causing cerebellar atrophy with postnatal onset do not lead to a significant reduction in pontine size, even in advanced stages. Variable cerebral involvement, including atrophy and delayed myelination, may be present.

The morphologic pattern of pontocerebellar hypoplasia is not specific for PCHs and has also been shown in other selected malformations, disruptions (eg, in extreme prematurity), and neurometabolic diseases (eg, in the congenital disorder known as glycosylation type 1A [OMIM 212065]) (48,49). Patients with mutations in CASK present with ataxia, nystagmus, postnatal microcephaly, severe cognitive impairment, and sensorineural hearing loss (50). The neuroimaging correlate includes severe global cerebellar hypoplasia, pontine hypoplasia, reduced gyral pattern, and unmyelinated corpus callosum (51,52). Mutations in CASK are inherited with an X-linked pattern. They are lethal in males and occur mostly in females. As mentioned earlier, PCH commonly involves the posterior fossa in migrational disorders and is associated with mutations in VLDLR and RELN in particular. Finally, PCH is a finding in congenital muscular dystrophies that are caused by defective dystroglycan O-glycosylation (discussed later).

**Joubert Syndrome.**—Joubert syndrome is characterized by hypotonia, ataxia, ocular motor apraxia, neonatal breathing dysregulation, and intellectual disability of variable severity (53,54). The “molar tooth sign” is the diagnostic criterion for Joubert syndrome and consists of elongated, thickened, and horizontally oriented superior cerebellar peduncles; a deep interpeduncular fossa; and vermian hypoplasia (Fig 8a, 8b) (55,56). Systemic involvement may be present and includes renal (nephronophthisis), ocular (colobomas, retinal dystrophy), hepatic (congenital hepatic fibrosis), and skeletal (various forms of polydactyly) involvement. Renal and hepatic involvement may cause high morbidity and mortality in patients with Joubert syndrome and necessitates appropriate work-up and regular follow-up. On the basis of the systemic involvement, six phenotypic subgroups have been described (57). Some degree of phenotype-genotype correlation has been shown between the six phenotypes and the 26 genes associated with Joubert syndrome thus far. All of these genes encode for proteins of the nonmotile primary cilia, which play a key role in the development and functioning of various cells, including retinal photoreceptors, epithelial cells lining the renal tubules and bile ducts, and neurons (54). In the central nervous system, the primary cilia are implicated in neuronal cell proliferation and axonal migration in the cerebellum and brainstem. In almost all patients, Joubert syndrome is inherited with an autosomal recessive pattern and poses a 25% recurrence risk in the affected family (54). Only mutations in OFD1 are exceptional and inherited with an X-linked pattern.

In Joubert syndrome, the spectrum of neuroimaging findings extends beyond the molar tooth sign and vermian hypoplasia and dys-
plasia, thereby confirming the heterogeneity of Joubert syndrome, not only from the clinical and genetic, but also from the neuroimaging point of view (56). The degree of vermian hypoplasia, form of the molar tooth sign, size of the posterior fossa, and shape and size of the cerebellar hemispheres are variable. Morphologic abnormalities of the brainstem are present in about 30% of patients and include a dysmorphic tectum and midbrain, thickening and elongation of the midbrain, and a small pons (56). Supratentorial involvement occurs in about 30% of patients and includes callosal dysgenesis, cephaloceles, hippocampal malrotation, migrational disorders, and ventriculomegaly (56). Differences in neuroimaging findings may be present in siblings, representing intrafamilial heterogeneity. No correlation between neuroimaging findings and genotype has been found; as a result, neuroimaging findings are of limited value in classifying patients with Joubert syndrome. Only the presence of a hypothalamic hamartoma allows differentiation between oral-facial-digital syndrome type VI and the other five Joubert syndrome subgroups (58).

Diffusion tensor imaging reveals the absence of decussation of the superior cerebellar peduncles and corticospinal tract, which implies an underlying defect in axonal guidance (Fig 8c) (59).

**Congenital Muscular Dystrophies Owing to Defective Dystroglycan O-Glycosylation.**—The “α-dystroglycanopathies” are a group of congenital muscular dystrophies resulting from mutations in 15 of the genes responsible for the O-glycosylation of α-dystroglycan. Mutations in these genes are inherited with an autosomal recessive pattern. The muscles, brain, and eyes are usually affected in α-dystroglycanopathies. Various phenotypes have been described based on the severity of the findings, including Fukuyama disease, muscle-eye-brain disease, and Walker-Warburg syndrome, which is the most severe form (60). However, isolated cerebellar involvement is possible (61).

Weakness, hypotonia, contractures, seizures, cognitive impairment, and eye involvement (eg,
Muscle-eye-brain disease and POMGnT1 mutation in a 5-month-old infant who presented with muscular hypotonia, weakness, intractable seizures, and poor visual function. (a) Sagittal T2-weighted MR image shows hypoplasia of the vermis, flattening of the ventral pons, a dysmorphic tectum and midbrain (arrow), an abnormal concave posterior border of the brainstem (*), an enlarged fourth ventricle, and supratentorial ventriculomegaly (arrowhead). (b) Coronal T2-weighted MR image shows cerebellar hypoplasia, multiple bilateral subcortical cysts in the cerebellar hemispheres (arrow), cerebellar dysplasia, generalized polymicrogyria (arrowheads), abnormal signal intensity of the supratentorial white matter, absence of the septum pellucidum, and marked ventriculomegaly.

Infratentorial involvement includes cerebellar hypoplasia or dysplasia, cerebellar cysts, pontine hypoplasia, ventral pontine cleft, and pontomesencephalic kinking (Fig 9) (60). Cerebellar cysts are a relatively specific finding. These cysts are most likely formed from the portions of the subarachnoid space that were engulfed by the dysplastic cerebellar folia, particularly at the boundary between normal and dysplastic cerebellar cortex (60). Supratentorial involvement ranges from mild ventriculomegaly, diffuse periventricular white matter changes, and focal areas of polymicrogyria (Fig 9b) to severe hydrocephalus, generalized white matter signal changes, and diffuse cortical abnormalities, including cobblestone lissencephaly (60,61).

Predominantly Brainstem Malformations

Pontine Tegmental Cap Dysplasia.—Pontine tegmental cap dysplasia is characterized by a flattened ventral pons, vaulted pontine tegmentum (the “cap”), partial absence of the middle cerebellar peduncles, vermian hypoplasia, a molar tooth–like aspect of the pontomesencephalic junction, and absent inferior olivary prominence (Fig 10a, 10b) (63).

Clinical findings are characterized by the involvement of vestibulocochlear, facial, trigeminal, and glossopharyngeal nerves, resulting in hearing loss, facial paralysis, trigeminal anesthesia, and difficulty in swallowing (63). Systemic involvement with vertebral segmentation anomalies, rib malformations, and congenital heart defects has been observed. The prognosis of pontine tegmental cap dysplasia appears to be highly variable, ranging from mild delay in cognitive development to severe disability (8,64,65). Pontine tegmental cap dysplasia is a sporadic malformation with unknown genotype and no familial recurrence.

Conventional neuroimaging findings are pathognomonic. Diffusion tensor imaging shows absence of the transverse pontine fibers and the presence of a dorsal transverse axonal band at the level of the so-called cap in the dorsal pons (Fig 10c) (63,64,66). The dorsal ectopic axonal band most likely results from abnormal axonal guidance and/or neuronal migration (66).

Horizontal Gaze Palsy with Progressive Scoliosis.—Horizontal gaze palsy with progressive scoliosis (OMIM 607313) is a rare autosomal recessive disorder caused by mutations in ROBO3, which encodes a receptor required for axonal guidance (67). Horizontal gaze palsy with progressive scoliosis is characterized clinically by congenital absence of horizontal eye movements, preservation of vertical gaze and convergence, and progressive development of scoliosis in childhood (68).

Neuroimaging reveals a butterfly-shaped medulla due to the missing prominence of the gracile...
and cuneate nuclei, as well as prominent inferior olivary nuclei with respect to the medullary pyramids (69). The pons is hypoplastic and has a dorsal midline cleft with absence of the bulging contour of facial colliculi (69). Diffusion tensor imaging typically shows the absence of decussation of the corticospinal tracts, pontine sensory tracts, and superior cerebellar peduncles (70).

Cerebellar Disruptions
Cerebellar maturation and adaptation follow a highly orchestrated series of programmed developmental processes of migration, proliferation, and arborization, starting in the middle of the 1st trimester and ending at about 2 years of age (72). The complexity and long duration of cerebellar development place the immature, developing cerebellum at high risk for acquired injury. The cerebellum is particularly vulnerable to metabolic and toxic insults as well as prenatal infections and hemorrhages, whereas it is less vulnerable to prenatal, perinatal, and postnatal hypoxic-ischemic events (5,73).

Predominantly Midbrain Malformations
Few disorders that selectively affect the midbrain have been reported. Zaki et al (71) recently reported a new malformation that is characterized by dysplasia of the diencephalic-mesencephalic junction with a butterfly-like contour of the midbrain on axial images.

Cerebellar Agenesis
Cerebellar agenesis is characterized by the near-complete absence of cerebellar tissue. The definition of cerebellar agenesis is based on the morphologic pattern and does not suggest the pathogenesis (5,74). Cerebellar agenesis may represent a malformation resulting from a genetically mediated pathomechanism (eg, mutations in PTF1A) or a disruption (eg, hemorrhage that occurs during gestation or in the perinatal period) (5,75).

All patients with cerebellar agenesis are symptomatic. Patients who survive infancy have variable degrees of cerebellar dysfunction (truncal and limb ataxia, dysarthria) as well as cognitive impairment (5). Neonates with cerebellar agenesis should be evaluated for diabetes mellitus, since this association is suggestive of a mutation in PTF1A (76).
Near-complete absence of cerebellar structures with remnants of the anterior vermis, floccules, and middle cerebellar peduncles are present at neuroimaging (Fig 11). In addition, a secondary pontine hypoplasia and a normal or enlarged posterior fossa may be seen.

**Global Cerebellar Hypoplasia**

Global cerebellar hypoplasia may result from several causes, both malformations and disruptions; hence, the widely varying clinical presentations and poor correlation with neuroimaging findings. With respect to malformations, global cerebellar hypoplasia has been reported in chromosomal abnormalities (eg, trisomies 13 and 18), metabolic disorders (eg, Zellweger or Smith–Lemli–Opitz syndrome), several genetic syndromes (eg, CHARGE syndrome or velocardiofacial syndrome), and migrational disorders (77). With respect to disruptions, global cerebellar hypoplasia may result from prenatal infections such as CMV or prenatal exposure to anticonvulsant drugs, alcohol, or cocaine (5).

Clinical features and prognosis are heterogeneous and reflect the specific cause. In terms of prognosis and recurrence risk, it is important to identify global cerebellar hypoplasia secondary to a malformation or disruption. This is usually possible on the basis of clinical history and findings as well as additional neuroimaging findings.

Neuroimaging findings show a cerebellum with normal or near-normal shape, but with a reduction in the volume and prominence of the subarachnoid spaces (Fig 12a). Involvement of the vermis and cerebellar hemispheres is typically similar. Ancillary supratentorial findings such as calcification, white matter signal abnormalities, and migrational disorders suggest a prenatal infection such as CMV (Fig 12b).

**Unilateral Cerebellar Hypoplasia**

Unilateral cerebellar hypoplasia is a rare finding with a spectrum of features ranging from complete aplasia to mild asymmetry in the size of the cerebellar hemispheres (5,78). Increasing experience with prenatal US and fetal MR imaging has demonstrated that unilateral cerebellar hypoplasia is of prenatal origin (and thus represents a disruption), with hemorrhage as the leading cause (5,79).

The most common clinical findings are delayed development and speech, hypotonia, ataxia, and abnormal ocular movements (80). The outcome is variable, ranging from near-normal development to marked motor and intellectual disability. Involvement of the cerebellar vermis is often, but not consistently, associated with a poorer cognitive outcome (78).

Unilateral cerebellar hypoplasia is characterized by variable involvement and volume loss in the cerebellar hemisphere and vermis, with a normal-sized posterior fossa (Fig 13). Evidence of hemorrhage may be present and is best visualized with susceptibility-weighted sequences (10). However, the absence of hemosiderin does not exclude hemorrhage, since the blood-brain barrier is permeable to hemosiderin-laden macrophages between 24–32 weeks gestation. Additional supratentorial disruptive lesions such as porencephalic cysts or clefts may be present and support the disruptive origin of unilateral cerebellar hypoplasia.
Cerebellar Cleft

Cerebellar cleft is a rare entity that primarily affects the cerebellar cortical gray matter, extends from the surface of the hemisphere into the parenchyma, and may reach the fourth ventricle, but without involvement of the vermis (81). Cerebellar clefts represent residual disruptive changes, most often secondary to fetal cerebellar hemorrhages (81). Supratentorial disruptive lesions such as schizencephaly may be present and thereby support the disruptive pathomechanism (81).

Clinical findings include language and speech disorders, truncal ataxia, ocular movement disorders, cognitive impairment, and behavioral disorders. The prognosis and outcome depend mostly on the presence and extent of supratentorial lesions.

Typical neuroimaging findings include (a) a cleft extending from the surface of the cerebellum through the cerebellar parenchyma toward the fourth ventricle, with malorientation of the cerebellar folia; (b) an irregular gray matter–white matter junction; and (c) abnormal arborization of the white matter confined to the region adjacent to the cleft, along with volume reduction in the affected hemisphere (Fig 14) (81).

Vanishing Cerebellum in Myelomeningocele

In a non–skin-covered myelomeningocele, prenatal hindbrain herniation through the foramen magnum is believed to be secondary to a prolonged prenatal CSF leak. This herniation may result in parenchymal damage due to ischemia, presumably induced by mechanical distortion, and marked reduction in the size of the cerebellar hemispheres (Fig 15). Boltshauser et al (82) reported this entity in three children with asymmetric involvement of the cerebellar hemispheres and sparing of the vermis.
Figure 15. Vanishing cerebellum in a 5-month-old male infant after surgical closure of an open myelomeningocele at birth. (a) Sagittal T2-weighted MR image shows a small posterior fossa, flattening of the ventral pons (white arrow), and small remnants of cerebellar tissue herniating into the spinal canal (white arrowhead). Typical findings of a Chiari 2 malformation, including an abnormal tectum (black arrowhead), a prominent massa intermedia (*), and dysgenesis of the corpus callosum (black arrows), are also seen. (b) Coronal T2-weighted MR image shows marked reduction in the size of the cerebellar hemispheres, with only small remnants being visualized (arrows).

Cerebellar Disruption or Injury Secondary to Prematurity

During the period from 28 weeks gestation to term, the surface area of the cerebellar cortex increases more than 30-fold (83). This rapid growth of the cerebellum follows a well-programmed developmental process and requires a high degree of energy, placing the developing cerebellum at high risk for injury, particularly during gestational weeks 24–32 (84). Arrested cerebellar development with reduction in the final cerebellar volume is the long-term sequela of cerebellar injuries in preterm neonates (83).

Cerebellar injury occurs in up to 20% of preterm infants prior to 32 weeks gestation (85). Impaired cerebellar development in preterm newborns may be caused by primarily destructive injuries or secondary lesions. Primarily destructive injuries include hemorrhage and ischemia and cause cerebellar tissue loss (86). Secondary lesions lead to underdevelopment of the cerebellum and are caused by direct (e.g., hemosiderin–blood products, glucocorticoid exposure, undernutrition) or indirect (impaired transsynaptic trophic effects) effects on the cerebellum (86). General perinatal risk factors for cerebellar injury in extremely premature neonates include prolonged ventilation, pressor support, posthemorrhagic hydrocephalus, and infratentorial hemosiderin deposition (84,87).
Surviving preterm neonates with cerebellar injuries have a high prevalence of motor, cognitive, and behavioral disabilities, including autistic features (86).

Typical neuroimaging findings of secondary cerebellar injury include volume reduction that predominantly involves the “skeletonized” cerebellar hemispheres compared with the less affected vermis; a flattened ventral pons; and an enlarged, balloon-shaped fourth ventricle (Fig 16) (49). Hemosiderin deposition may be present and is best depicted with susceptibility-weighted sequences.

**Conclusion**

Congenital posterior fossa abnormalities represent a wide variety of disorders that include both malformations and disruptions. These disorders may involve a single structure within the posterior fossa (eg, the cerebellum alone) or a combination of structures (eg, the pons and cerebellum). Neuroimaging yields detailed anatomic findings and plays a key role in the diagnosis of congenital posterior fossa abnormalities. Well-defined neuroimaging-based diagnostic criteria are available for the various disorders and should be used by pediatric neurologists, general radiologists, and neuroradiologists. Accurate classification of congenital posterior fossa abnormalities is important for therapy, prognosis, and genetic counseling. If the diagnostic criteria for well-defined anomalies are not met, a detailed anatomic description is to be preferred, and confusing terms that lack specificity (eg, Dandy-Walker variant, Dandy-Walker spectrum) should be avoided or, even better, abandoned.

**References**


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