**Bilateral Posterior Periventricular Nodular Heterotopia: A Recognizable Cortical Malformation with a Spectrum of Associated Brain Abnormalities**

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**ABSTRACT**

**BACKGROUND AND PURPOSE:** Bilateral posterior PNH is a distinctive complex malformation with imaging features distinguishing it from classic bilateral PNH associated with FLNA mutations. The purpose of this study was to define the imaging features of posterior bilateral periventricular nodular heterotopia and to determine whether associated brain malformations suggest specific subcategories.

**MATERIALS AND METHODS:** We identified a cohort of 50 patients (31 females; mean age, 13 years) with bilateral posterior PNH and systematically reviewed and documented associated MR imaging abnormalities. Patients were negative for mutations of FLNA.

**RESULTS:** Nodules were often noncontiguous (n = 28) and asymmetric (n = 31). All except 1 patient showed associated developmental brain abnormalities involving a spectrum of posterior structures. A range of posterior fossa abnormalities affected the cerebellum, including cerebellar malformations and posterior fossa cysts (n = 38). Corpus callosum abnormalities (n = 40) ranged from mild dysplasia to agenesis. Posterior white matter volume was decreased (n = 22), and colpocephaly was frequent (n = 26). Most (n = 40) had associated cortical abnormalities ranging from minor to major (polymicrogyria), typically located in the cortex overlying the PNH. Abnormal Sylvian fissure morphology was common (n = 27), and hippocampal abnormalities were frequent (n = 37). Four family cases were identified—2 with concordant malformation patterns and 2 with discordant malformation patterns.

**CONCLUSIONS:** The associations of bilateral posterior PNH encompass a range of abnormalities involving brain structures inferior to the Sylvian fissures. We were unable to identify specific subgroups and therefore conceptualize bilateral posterior PNH as a continuum of infrasylvian malformations involving the posterior cerebral and hindbrain structures.

**ABBREVIATIONS:** DHPLC = denaturing high-performance liquid chromatography; FLNA = Filamin A gene; PNH = periventricular nodular heterotopia

Subependymal or periventricular nodular heterotopia is the most commonly identified gray matter heterotopia in clinical practice.\(^1\)\(^-\)\(^5\) It may be more often associated with other malformations than commonly appreciated.\(^5\) PNH is a collection of subependymal nodules of gray matter found along the walls of the lateral ventricles.\(^1\)\(^-\)\(^9\) PNH consists of clusters of neurons that fail to migrate away from the embryonic ventricular zone to the developing cerebral cortex. Radial migration occurs maximally between the seventh and sixteenth gestational weeks, along radial glial fibers.\(^10\)\(^-\)\(^13\) The major locus associated with PNH is the *Filamin A* gene on Xq28 that causes X-linked PNH that is typically bilateral and symmetric. This classic form occurs in females, many of whom have normal intelligence and epilepsy.\(^14\) The nodules in these patients involve predominantly the anterior bodies and frontal horns of the lateral ventricles and are often diffuse, contiguous, and symmetric.\(^9\)\(^,\)\(^15\)\(^-\)\(^17\)

Our study was motivated by the observation that the second most common pattern of PNH showed nodules predominantly along the atria and temporal horns of the lateral ventricles. Patients with this posterior form of PNH were *FLNA*-negative, had symptoms that appeared different from those in the classic form, and were a sizeable group that were underappreciated as a separate entity in the literature. Reported subtypes of *FLNA*-negative
PNH include PNH with hydrocephalus, PNH with microcephaly; PNH with frontonasal dysplasia; PNH with limb abnormalities; PNH with fragile X syndrome; PNH with ambiguous genitalia; and, micronodular, unilateral, laminar, ribbonlike, and linear PNH. We also observed that patients with posterior PNH had more frequent abnormalities of other brain structures compared with patients with classic bilateral PNH. We therefore questioned whether bilateral posterior PNH is a distinct entity compared with classic PNH and reviewed all cases in which the nodules were primarily posterior and bilateral. We examined the associated brain abnormalities to determine whether there were subcategories within the posterior PNH group or whether posterior PNH is part of a spectrum of disordered infra-Sylvian development.

MATERIALS AND METHODS

Imaging

We identified 50 patients with posterior-predominant PNH on MR imaging scans referred by multiple centers. Structural MR imaging examinations were performed between 1996 and 2009. All patients had multiplanar multiecho sequences, including T1, T2, and FLAIR. T1 images were of high quality, typically by using an inversion-prepared sequence, and were of appropriate section thickness, typically 3 mm. T1- and T2-weighted images were available in all cases.

We classified each patient’s MR images on the basis of imaging findings. The shape, location, and symmetry of PNH were analyzed, and the presence of all associated abnormalities was documented. Apart from describing the features of the PNH, we assessed associated brain structures as follows:

1. Cerebellum: morphology of the vermis and hemispheres
2. Posterior fossa: presence or absence of a posterior fossa cyst
3. Brain stem morphology
4. Corpus callosum: the presence and shape of the corpus callosum
5. White matter: volume and signal intensity
6. Ventricles: morphology and size of the ventricles. Lateral ventricles were assessed for colpocephaly, defined as disproportionate dilation of the trigones and occipital horns of the lateral ventricles.
7. Hippocampi: symmetry, shape, and signal intensity of the hippocampi. Hippocampi were assessed as asymmetric if the shape or size of the 2 sides differed but the internal architecture was preserved with normal signal intensity and normal hippocampal rotation. Dysplasia was assessed by using the following criteria: shape, size, signal intensity, and hippocampal rotation.
8. Cerebral cortex: The cortical mantle was assessed for the presence of cortical malformations such as cortical dysplasia and polymicrogyria.

Filamin A Analysis

FLNA was examined by either DHPLC (Transgenomic, Omaha, Nebraska) or Sanger sequencing by using BigDye ter-

RESULTS

Patient Characteristics

The 50 patients ranged in age from a 26-week fetus (in utero scan) to 67 years of age, with a mean postnatal age of 13 years and a median age of 6 years at the time of the first MR imaging. There were 31 females (62%). Forty-two were unrelated individuals. There were 2 sister pairs, 1 monozygotic twin brother pair, and 1 mother-son pair. The son was imaged as a fetus with 2 fetal MR imaging scans at 26 weeks’ and 32 weeks’ gestation as well as with a postnatal study.

Nodules

We defined posterior PNH as either restricted to or predominant in the region around the posterior horns, atria, or temporal horns of the lateral ventricles. Heterotopic nodules had an exclusively posterior infrasylvian distribution in 35 (70%) patients, while 15 (30%) had a mixed picture with a predominance of posterior nodules and scattered nodules anteriorly along the bodies of anterior horns. Nodules were contiguous in 22 (44%) patients and symmetric in 19 (38%) patients.
Posterior Fossa Abnormalities

Thirty-eight of 50 (76%) patients had a range of posterior fossa abnormalities (Fig 1). The major abnormalities included severe dysplasia/hypoplasia of the cerebellar hemispheres with large CSF fluid collections in the Dandy-Walker continuum. Twenty of 50 (40%) patients had dysplastic cerebella with vermian abnormalities. Seventeen patients with cerebellar abnormalities had posterior fossa cysts. The appearance of the cysts varied from small unilateral cysts with no mass effect to large cysts expanding the posterior fossa. The latter were associated with extremely small and malformed cerebellar hemispheres. Minor abnormalities included deficiency of the tentorium cerebelli. Five patients had small but morphologically normal cerebella with posterior fossa cysts (Fig 2).

With 2 exceptions, all patients with cerebellar abnormalities also had abnormalities of the corpus callosum and were more likely to have white matter abnormalities. Ten patients (20%) had brain stem abnormalities, including a patient with a tectal plate glioma and 6 patients with tectal plate thickening. The other 3 had unilateral or bilateral brain stem hypoplasia.

Corpus Callosum and White Matter

Forty of 50 (80%) patients demonstrated abnormalities of the corpus callosum ranging from mild dysplasia to complete agenesis. The corpus callosum abnormalities were divided into 1 of 4 major groups: agenesis (6 patients), thin (21 patients), thick (4 patients), and minor irregularities (4 patients), (Figs 3 and 4).

All patients had abnormalities of the whole or the posterior half of the corpus callosum. There were 3 patients with complete agenesis of the corpus callosum and 3 with partial agenesis who had absence of the posterior body and splenium. Five of these patients had abnormal hippocampi. Twenty-three (46%) patients had thin posterior bodies and splenia of the corpus callosum with normal anterior components, while 3 patients had thinning of the entire corpus callosum. Four patients had an unusual thickened appearance of the corpus callosum with dysplastic splenia and no colpocephaly. All 4 had associated hippocampal dysplasia and abnormal Sylvian fissures.

Fifteen patients with callosal abnormalities had a normal cerebellum, while 7 patients had a normal cerebellum and a normal corpus callosum. These patients were more likely to have hippocampal abnormalities and cortical/sulcation abnormalities, ranging from Sylvian fissure abnormalities to areas of cortical thickening and suspected transmantle bands.

Twenty-two (44%) patients demonstrated decreased posterior white matter volume, and 5 patients had nonspecific increased T2 signal intensity within the white matter. Two of these had unilateral anterior temporal white matter signal abnormality of the type seen in temporal lobe dysplasias or hippocampal sclerosis. One patient demonstrated delayed myelination for age, and 1 had white matter distortion due to multiple nodules.

Ventricles

Thirty-two (64%) patients had ventricular abnormalities. Of these, 26 patients had colpocephaly. In 3 cases, the ventricle outlines were very distorted by the presence of the nodules. There was mild generalized ventriculomegaly in 3 patients.
Forty (80%) patients had associated cortical mantle or sulcation abnormalities, usually in the cortex directly overlying the PNH. These included nonspecific cortical thickening, linear transmantle bands, cortical dysplasia, and polymicrogyria (On-line Table 1). Linear transmantle bands were T1 hypointense and T2 hyperintense lines extending from the nodules to the overlying cortical surface. Twelve of 50 (24%) had cortical dysplasia, schizencephaly, or polymicrogyria. There were 11 patients with sulcation abnormalities alone; 4 with transmantle bands alone; and 13 with combinations of cortical thickening, linear transmantle bands, and sulcation abnormalities. Twenty-seven (54%) patients had abnormal Sylvian fissures with a spectrum of appearances as shown in Fig 5.

**Hippocampal Abnormalities**

There was a spectrum of hippocampal abnormalities in 37/50 (74%) patients. Twenty-five patients had dysplastic hippocampi, of which 16 were also malrotated. Twelve (24%) had asymmetric hippocampi (Fig 6).

**Additional Abnormalities**

Five patients had abnormalities of the pituitary gland. These included an absent posterior pituitary hyperintense neurohypophysis ("bright spot") \((n = 1)\), an ectopic posterior pituitary hyperintense neurohypophysis \((n = 2)\), a pituitary cyst \((n = 1)\), and a duplicated pituitary hyperintense region \((n = 1)\). One patient had an occipital encephalocele repaired as an infant with associated venous anomalies. One patient had lobar holoprosencephaly. A pair of female siblings had congenital cataracts with optic nerve hypoplasia.

**Familial Cases**

There were 4 families with 2 affected individuals in each family (2 sibling pairs, 1 pair of monozygotic twins, and a mother-son pair). The first sister pair and the mother-son pair showed diverse malformations, while the twins and the second sister pair had concordant malformations (Fig 7 and On-line Tables 1 and 2).

**Filamin A Analysis**

Examination of all coding exons and intron-exon boundaries of FLNA was completed for 40 subjects. No point mutations or small insertion/deletions confined to a single exon were detected in any subject. Additionally, no deletions or duplications of either the entire locus or involving the assayed exons were detected. Thus there were no sequencing mutations or copy number variants of FLNA found in the tested cohort with bilateral posterior PNH.

**DISCUSSION**

Bilateral posterior PNH is a cortical malformation entity showing a spectrum of disordered brain development and is distinct from classic bilateral PNH due to not only the location of the nodules but also to the common occurrence of malformations of other brain structures and the absence of mutations in the FLNA gene. We found only 1 case that had no identifiable malformations other than the PNH. This is despite the selection criteria being only bilateral posterior nodules, admittedly in the context of a tertiary referral center. In some cases these abnormalities were only detected in this systematic review and were not reported on the routine studies that led to recruitment. We suggest, therefore, that posterior bilateral PNH is typically associated with other brain malformations and that it is part of a more diffuse process of posterior or infrasylvian brain maldevelopment other than just a disorder of neuronal migration.
Associated Malformations

Cortical abnormalities occurred in 80% of cases, usually in the cortex overlying the PNH. Sylvian fissures were often abnormal (50%), some demonstrating minor anomalies such as elongation and some with severe abnormalities such as bilateral perisylvian polymicrogyria. These could be secondary to a disturbance in a range of fundamental forces shaping the cortex, including processes underlying the normal gyrification of the brain, such as genetic control of sulcation, cytoarchitecture, differential cortical growth, and white matter tension morphogenesis.21-28

Posterior callosal abnormalities were found in 80% of patients. This was typically thinning or agenesis and likely reflects disordered or failed migration or defective axon guidance. Although the recognized association of callosal and hippocampal abnormalities was seen in our patients, we did not see this combination as a separate subgroup.1,3,29-31 The arrested hippocampal rotation is likely to be due to secondary effects of abnormal development, though we acknowledge it could be part of the same genetic defect that arrests the migration of neural progenitor cells.32 Four patients had callosal hypertrophy with dysplastic splenia, hippocampal dysplasia, and abnormal Sylvian fissures. A hyperplastic corpus callosum is rare and has been reported in neurofibromatosis type 133 and Cohen syndrome34,35 and likely reflects a failure of normal pruning of the commissural fibers.36,37

We commonly saw posterior ventricular dilation (50%), which we suspect has sometimes been interpreted as hydrocephalus.9 Our interpretation is that this is “colpocephaly,” because our cases had no clinical or other imaging features suggesting raised intracranial pressure.

Cerebellar abnormalities of varying severity were found in 50% of patients and varied from asymmetric cerebellar hemisphere hypoplasia, universally associated with a small or absent vermis, to a “microcerebellum” that was small but normally formed. A bulky vermis was seen in 5 patients, likely due to nodular heterotopia causing expansion. One patient had a small tectal plate lesion that mimicked gray matter on all sequences and did not enhance, suggesting a periventricular nodule rather than an indolent midbrain tumor.38 Previously reported pituitary gland abnormalities were seen in 10% of patients, including one with an ectopic posterior pituitary gland.39

Genetics of the Bilateral Posterior PNH Spectrum

At least some bilateral infra-Sylvian PNH has a genetic basis as demonstrated by our 4 kinships that include identical twins and a mother-son pair. Each related pair of individuals showed a range of abnormalities within the bilateral posterior PNH “spectrum.” This is strong clinical evidence of a genetic basis underlying critical aspects of this developmental disorder. The twins and second sister pair were concordant with respect to associated malformations within each pair. The first sister pair and the mother-son pair showed variation in their malformations. There were no clear environmental or acquired factors to explain the disparity in the related pairs. These differences with presumed common genes, best illustrated by the mother-son pair, indicate that there may be a spectrum of expressions of common mutations or modifying factors. This suggests that common pathways may be involved in the range of features we see in this bilateral PNH cohort.

Bilateral posterior PNH is a distinct malformation of cortical development associated with a spectrum of infra-Sylvian developmental abnormalities. The fact that we could identify a series of 50 such cases suggests that posterior PNH is a large subgroup of the common bilateral PNH cases. In our experience of ~200 patients with bilateral PNH, it is the second
In nonsyndromic bilateral posterior PNH, Parriini et al. distinguished 3 major associations: hippocampal malformation and cerebellar hypoplasia (n = 10), polymicrogyria (n = 2), and hydrocephalus (n = 5). We selected patients who had similar PNH features but could not easily fit them into these 3 specific categories. Rather than discrete subgroups, we suggest that bilateral posterior PNH is a distinct malformation entity that encompasses a continuum of brain malformations involving posterior or infrasylvian structures (Fig 8).

**CONCLUSIONS**

While periventricular nodules have been shown in pathologic studies to have mostly sparse connections with extranodular regions and with other adjacent nodules, some nodules have been shown to be more connected to overlying and potentially abnormal cortex than others. 

Epileptic activity may be due to a combination of the intrinsic epileptogenicity of the nodule as well as its connections to cortex.

**FIG 8.** Summary of the malformation spectrum seen in bilateral posterior PNH, highlighting the infrasylvian location of the nodules and the associated malformations. Black nodules represent typical frontocentral nodules and blue nodules represent infrasylvian location.

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