Periventricular Heterotopia: New Insights into Ehlers-Danlos Syndrome

Volney L. Sheen, MD, PhD and Christopher A. Walsh, MD, PhD

Nature often employs similar mechanisms to complete similar tasks, thus the evolution of homologous proteins across various organ systems to perform similar but slightly different functions. In this respect, disorders attributed to specific genetic mutations, while initially thought to be restricted in function and purpose, may provide broad insight into general cellular and molecular mechanisms of development and maintenance. One such example can be seen in the brain malformation, periventricular heterotopia (PH), which is characterized by very specific nodules of neurons that line the lateral ventricles beneath the cerebral cortex. PH is seen as a disorder of neuronal migration and can be caused by mutations in filamin A (FLNA), which encodes an actin-binding protein that regulates the cytoskeleton and cell motility. Recent advances in our understanding of the genetic causes of PH suggest that mutations in this gene, however, are also associated with the connective tissue disorder, Ehlers-Danlos syndrome (EDS), in which affected individuals present with joint and skin hyperextensibility and vascular problems including aortic dissection, excessive bleeding and bruising. While much still remains unknown regarding the mechanistic role of FLNA in giving rise to PH and EDS, a common cellular and molecular basis likely gives rise to these two seemingly unrelated clinical disorders.

Keywords: Central nervous system/pathology; Collagen; Connective tissue; Cortical development; Ehlers-Danlos syndrome; Filamin A; Inherited disorders, CNS; Periventricular heterotopia

Genetics of Periventricular Heterotopia and Ehlers-Danlos Syndrome

Periventricular heterotopia (PH) has been interpreted as a disorder in neuronal migration where newly born neurons fail to migrate from their birthplace along the lining of the lateral ventricles of the brain, thereby giving rise to the characteristic ectopically placed neuronal nodules (figure 1). Mutations in two genes have been identified as the genetic cause of PH. The more common X-linked dominant form of PH is due to mutations in the filamin A (FLNA) gene.1,2 At least 80% of all familial cases of PH are due to a FLNA mutation with some 20% of sporadic cases attributable to this gene.3 A much more rare autosomal recessive form of PH with microcephaly results from mutations in the ARFGEF2 gene which encodes the brefeldin-inhibited guanine exchange factor 2 (BIG2) protein.4 Filamin A interacts with the actin cytoskeleton and thereby regulates various aspects of cell shape, motility and function.5 BIG2 regulates vesicle trafficking from the trans-Golgi apparatus to the cell surface in various tissues of the body. Potentially, the secreted or surface proteins transported through BIG2-dependent vesicles may interact with filamin A near the cell surface and disruption of these as proteins could give rise to PH by impairing neuronal migration.

Ehlers-Danlos syndrome (EDS) describes a heterogeneous group of disorders, which mainly affect the skin, joints and vasculature, but uniformly involve defects in collagen. Diagnosis of EDS and sub-classification into one of the six major subtypes of EDS is based on major (skin hyperextensibility, joint hypermobility, skin fragility and scarring, vascular fragility, kyphoscoliosis and scleral involvement) and minor diagnostic criteria.6 Mutations in different collagens and collagen-modifying enzymes have been recognized for all sub-types of EDS with the exception of type III hypermobility-type EDS.7 The collagens represent a large family of structurally related extracellular matrix proteins essential for development, cell attachment, platelet aggregation and for providing tensile strength to the
EDS. While affected patients have nodular brain with PH who have independently been identified as having cases of PH and EDS (8 females and 1 male), three presentation of EDS which suggests that the clinical features clear correlation exists between the type of mutation and resulting in a premature stop Q1248X in this family has subsequently detected a C3742T base pair change to chromosome Xq28, the hemizygous male lethality. Initial studies suggested linkage such as EDS follows that expected of an X-linked dominant disorder, including joint hypermobility, skin fragility and aortic heterotopia, they also exhibit features consistent with EDS, gene.2 Denaturing high performance liquid chromatography Western blot analyses suggesting some disruption in this mutation, but filamin A protein could not be detected on EDS, one pedigree was initially reported as having no exonic single base pair deletion). Of the two families with PH and deletion, C116G resulting in an A39G change, 4147 del G mutations were identified (2762 del G single base pair). Of the nine sporadic mutations in patients FLNA mutations. Approximately 90% of patients have focal epilepsy.14 Retroactive studies would suggest that the prognostic outcome of the epilepsy is dependent upon the electrographic and neuroradiological findings for the particular patient with PH. When imaging demonstrates PH in the absence of other brain malformations, patients generally have normal intelligence with their seizures beginning in the second decade of life. The EEG often shows only focal abnormalities, and the seizures usually do not become frequent and generally disappear. When the PH is associated with other cortical or cerebral malformations, patients may have mental retardation, and the seizures generally begin during the first decade of life. The EEG often shows focal and bitemporal abnormalities, and the seizures usually do not begin in the second decade of life. The EEG often shows focal and bitemporal abnormalities, and the seizures usually do not begin in the second decade of life. Neurocognitive studies, however, suggest that some 80% of individuals with PH harbor FLNA mutations.1,10 Several recurrent missense mutations in the actin-binding and rod domains of the FLNA gene have likewise been implicated in malformations affecting craniofacial structures and the skeleton.11-13 These syndromes, however, are thought to arise from a gain-of-function mutation in this gene. Overall, the broad spectrum of abnormalities attributable to FLNA mutations mirrors the widespread abnormalities in the vasculature, gut and bone associated with connective tissue disorders.

Many features of EDS can also be seen in affected individuals with PH but no formal diagnosis of a connective tissue disorder. Patients have been reported to have patent ductus arteriosus, bicuspid aortic valve, or aneurysmal dilatation of the sinus of Valsalva and/or thoracic aorta at young ages, but the true prevalence of these cardiac abnormalities are not yet known.1 Hemizygous males who lack a normal copy of the X-linked FLNA gene generally die prenatally from excessive bleeding.9 Congenital gut and genitourinary anomalies have also been reported in boys harboring FLNA mutations.1,10 Several recurrent missense mutations in the actin-binding and rod domains of the FLNA gene have likewise been implicated in malformations affecting craniofacial structures and the skeleton.11-13 These syndromes, however, are thought to arise from a gain-of-function mutation in this gene. Overall, the broad spectrum of abnormalities attributable to FLNA mutations mirrors the widespread abnormalities in the vasculature, gut and bone associated with connective tissue disorders.

Neurological symptoms (primarily seizures and dyslexia) are the most common presenting features associated with PH due to FLNA mutations. Approximately 90% of patients have focal epilepsy.14 retrospective studies would suggest that the prognostic outcome of the epilepsy is dependent upon the electrographic and neuroradiological findings for the particular patient with PH. When imaging demonstrates PH in the absence of other brain malformations, patients generally have normal intelligence with their seizures generally begin during the first decade of life. The EEG often shows focal and bitemporal abnormalities, and the seizures generally begin during the first decade of life. The EEG often shows focal and bitemporal abnormalities, and the epilepsy generally increases in frequency and may become refractory.15 Recent studies have also suggested a high incidence of dyslexia in the PH population.16 Since most patients with PH have normal intelligence, the dyslexia is not commonly recognized. Neurocognitive studies, however, suggest that some 80% of individuals with PH have deficits in reading, processing speed and executive function. The extent of this impairment is not correlated with the type of seizure medications or the severity of seizures, but is more severe when more widely distributed heterotopia is noted on imaging.

Potential Pathogenic Mechanisms in PH and EDS
The recent association between PH and EDS potentially broadens the current understanding of the molecular pathogenesis of EDS. EDS is typically viewed as a disorder of the extracellular matrix involving disruption in glycoproteins, such as collagen or proteoglycans such as

Figure 1. Radiographic phenotype of periventricular heterotopia. Axial T1-weighted magnetic resonance image of the brain demonstrates bilateral periventricular nodular heterotopia (white arrowheads) along the lateral ventricles. While not commonly appreciated, this particular patient also had small vessel ischemic changes at an early age, suggestive of some abnormality in the blood vessels.

connective tissues in bone, skin, ligaments and tendons, and blood vessels.8 Loss of the elasticity and strength of collagen in EDS presumably contributes to the structural failure in connective tissue of the various affected organ systems.

Clinical Presentations Associated with PH and FLNA Mutations
Recently, we have identified FLNA mutations in patients with PH who have independently been identified as having EDS.2 While affected patients have nodular brain heterotopia, they also exhibit features consistent with EDS, including joint hypermobility, skin fragility and aortic dilation in early adulthood. The inheritance pattern in PH and EDS follows that expected of an X-linked dominant disorder, such as FLNA with affected females and presumptive hemizygous male lethality. Initial studies suggested linkage to chromosome Xq28, the FLNA locus. Of the nine sporadic cases of PH and EDS (8 females and 1 male), three FLNA mutations were identified (2762 del G single base pair deletion, C116G resulting in an A39G change, 4147 del G single base pair deletion). Of the two families with PH and EDS, one pedigree was initially reported as having no exonic mutation, but filamin A protein could not be detected on Western blot analyses suggesting some disruption in this gene.2 Denaturing high performance liquid chromatography has subsequently detected a C3742T base pair change resulting in a premature stop Q1248X in this family (personal communications, Dr. Stephen Robertson). No clear correlation exists between the type of mutation and presentation of EDS which suggests that the clinical features of EDS may reflect some degree of variable penetrance.3 Moreover, it is not known whether individuals with EDS in the absence of PH may harbor FLNA mutations.
elastin or other extracellular matrix proteins including tenascin-X. Cells attach to the extracellular matrix by means of transmembrane glycoproteins called integrins, whose intracellular portion binds to the actin filaments of the cytoskeleton. Moreover, filamin A has been shown to bind the \( \alpha \beta \)-integrin cell adhesion receptors, but does not alter the formation of these focal adhesion sites and does not alter the extracellular matrix assembly. Thus, the finding of PH and EDS due to \( FLNA \) mutations would suggest a novel intracellular cause of EDS. Perhaps \( FLNA \) mutations cause EDS by decreasing the strength of attachment of the cells to the extracellular matrix or by impaired regulation of the cytoskeleton through the filamin binding to integrin proteins. Reflecting back on the brain defect, PH may not represent a disorder of cell motility. Rather, impaired structural integrity of the neuroependyma may weaken the ventricular lining during neurogenesis and lead to PH formation, much in the same manner by which the aorta dilates and the skin becomes overly extended from increased stress in EDS.

A more direct relationship between filamin and collagen can be appreciated when examining the mechanisms involved in platelet aggregation in which mutations in the \( FLNA \) (EDS variant of PH) and in \( COL3A1 \) (vascular type EDS) are both known to cause excessive bleeding, presumably from vascular fragility and platelet dysfunction. When the integrity of the vascular wall is compromised, as in vascular EDS, platelets adhere to subendothelial collagens exposed at the sites of blood vessel injury via the glycoprotein (GP) Ib-V-IX complex and integrin \( \alpha 2 \beta 1 \). This initial platelet entrapment requires the plasma protein von Willebrand factor, which simultaneously binds the GP and integrin complex and facilitates platelet tethering and rolling. More stable binding of collagen to the platelet collagen receptors (namely integrin \( \alpha 2 \beta 1 \) and GPVI) follows this initial interaction. In the vascular type of EDS, impaired collagen integrity disrupts this process. Likewise, filamin A binds both the integrin and the GPIb-IX complex, and studies suggest that the association of this receptor complex with the membrane skeleton, while not essential for platelet tethering or rolling under low shear conditions, is critical for maintaining platelet adhesion at high shear rates. Thus, the interactions between the extracellular collagen matrix, the platelet collagen receptors, filamin and the cytoskeleton collectively mediate platelet adhesion, activation and subsequent hemostasis.

Table 1. Widespread and overlapping features between Ehlers-Danlos syndrome and filamin related disorders.

<table>
<thead>
<tr>
<th>Types</th>
<th>Protein</th>
<th>Gene</th>
<th>Organ system</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ehlers-Danlos Syndrome</td>
<td>Type V procollagen</td>
<td>COL5A1/COL5A2</td>
<td>Skin hyperextensibility</td>
</tr>
<tr>
<td>Type I/II classic</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Joint hypermobility</td>
</tr>
<tr>
<td>Type III hypermobility</td>
<td>Type III procollagen</td>
<td>COL3A1</td>
<td>Blood vessel fragility</td>
</tr>
<tr>
<td>Type IV vascular</td>
<td>Lysyl hydroxylase 1</td>
<td>PLOD1</td>
<td>Muscular hypotonia</td>
</tr>
<tr>
<td>Type VI kypchoscoliotic</td>
<td>Type I procollagen</td>
<td>COL1A1/COL1A2</td>
<td>Joint hypermobility</td>
</tr>
<tr>
<td>Type VIIA/B arthrocralasia</td>
<td>Procollagen-N-proteinase</td>
<td>ADAMTS-2</td>
<td>Skin hyperextensibility</td>
</tr>
<tr>
<td>Type VIIIC dermatosparaxis</td>
<td>FEP1</td>
<td>FENL</td>
<td>Bone anomalies</td>
</tr>
</tbody>
</table>

Filamin Protein Syndromes

<table>
<thead>
<tr>
<th>Syndrome</th>
<th>Protein</th>
<th>Gene</th>
<th>Organ system</th>
</tr>
</thead>
<tbody>
<tr>
<td>Periventricular heterotopia (300049)</td>
<td>Filamin A</td>
<td>FLNA</td>
<td>Brain heterotopia</td>
</tr>
<tr>
<td>Otopalatodigital syndrome type I (311300)</td>
<td>Filamin A</td>
<td>FLNA</td>
<td>Bone anomalies</td>
</tr>
<tr>
<td>Otopalatodigital syndrome type II (304120)</td>
<td>Filamin A</td>
<td>FLNA</td>
<td>Bone anomalies</td>
</tr>
<tr>
<td>Frontometaphyseal dysplasia (305620)</td>
<td>Filamin A</td>
<td>FLNA</td>
<td>Bone anomalies</td>
</tr>
<tr>
<td>Melnick-Needles syndrome (309350)</td>
<td>Filamin A</td>
<td>FLNA</td>
<td>Bone anomalies</td>
</tr>
<tr>
<td>Ehlers-Danlos variant of periventricular heterotopia (300537)</td>
<td>Filamin A</td>
<td>FLNA</td>
<td>Joint hypermobility</td>
</tr>
<tr>
<td>Spondylocarpotarsal syndrome (272460)</td>
<td>Filamin B</td>
<td>FLNB</td>
<td>Bone anomalies</td>
</tr>
<tr>
<td>Autosomal dominant Larsen syndrome (150250)</td>
<td>Filamin B</td>
<td>FLNB</td>
<td>Joint hypermobility</td>
</tr>
<tr>
<td>Type I atelosteogenesis (108720)</td>
<td>Filamin B</td>
<td>FLNB</td>
<td>Bone anomalies</td>
</tr>
<tr>
<td>Type III atelosteogenesis (108721)</td>
<td>Filamin B</td>
<td>FLNB</td>
<td>Bone anomalies</td>
</tr>
<tr>
<td>Myofibrillar myopathy</td>
<td>Filamin C</td>
<td>FLNC</td>
<td>Muscle</td>
</tr>
</tbody>
</table>
the multi-organ system involvement in EDS, the broad distribution of filamin proteins reflects its widespread function and potential interactions with the extracellular matrix (see table 1). The filamin family of actin-binding proteins are composed of three highly homologous members: filamin A, filamin B and filamin C. Mutations in FLNA primarily involve abnormalities of the central nervous system, skeletal and vascular system.1,3 Mutations in the filamin B gene lead to skeletal and joint deformities.21 Mutations in the filamin C gene cause a myopathy.22

Tests Available in the Diagnosis of PH
In most cases, patients with PH have normal intelligence and no focal neurological findings on examination. Seizures are the most common presenting feature and the heterotopic nodules are incidentally observed following radiographic imaging.

Definitive diagnosis of PH is made through radiographic imaging. In X-linked PH due to FLNA mutations, magnetic resonance images uniformly demonstrate bilateral near-continuous nodular PH, typically in females. Findings of a thinned corpus callosum and mega cisterna magna are often associated with this disorder.1 There are no consistent radiographic differences between patients carrying different mutations.23 PH and mega cisterna magna, while subtle, can also be appreciated by ultrasonography during the second trimester.24 Positron emission tomography and single photon emission computed tomography scans suggest the glucose metabolism and perfusion of the heterotopic gray matter are identical to the normal cerebral cortex.25 In the autosomal recessive form of PH due to ARFGEF2 mutations, magnetic resonance images also demonstrate bilateral near-continuous nodular PH but have distinguishing features of microcephaly, mild ventriculomegaly and white matter changes.4

Definitive diagnosis of genetic causes of PH requires molecular testing for the genes, although this service is primarily performed on a research basis only.

Treatment of PH
No uniform guidelines exist for the management of complications from PH (namely seizures), although treatment generally follows standard guidelines involving focal epilepsies. In most cases, management through anti-convulsant medications is sufficient in controlling events.15 For intractable focal seizures requiring surgical resection, the best predictor of surgical outcome is the capacity to localize the focal epileptic generator, which may or may not include the heterotopia, through invasive recording by stereo-encephalography.26 Resection of the epileptogenic zones from patients with unilateral rather than bilateral PH also faired better.27

Conclusions
The diagnosis of PH and an awareness of potential problems associated with this disorder, particularly the risk of vascular dissection and dyslexia, should prompt a more careful initial evaluation and routine followup. An argument can be made for imaging of large vessels of the body given the risk of aortic dilation and rupture before the age of 40. Similarly, a battery of neuropsychological testing should be considered in instances where learning deficits are suspected.

References


Author Affiliations
Volney L. Sheen, MD, PhD
Department of Neurology
Beth Israel Deaconess Medical Center
Boston, Massachusetts

Christopher A. Walsh, MD, PhD
Division of Neurogenetics and Howard Hughes Medical Institute
Department of Neurology
Beth Israel Deaconess Medical Center
Boston, Massachusetts