A review of extraaxial developmental venous anomalies of the brain involving dural venous flow or sinuses: persistent embryonic sinuses, sinus pericranii, venous varices or aneurysmal malformations, and enlarged emissary veins

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This paper is a narrative review of extraaxial developmental venous anomalies (eDVAs) of the brain involving dural venous flow or sinuses: persistent embryonic sinuses, sinus pericranii, enlarged emissary veins, and venous varices or aneurysmal malformations. The article highlights the natural history, anatomy, embryology, imaging, clinical implications, and neurosurgical significance of these lesions, which the authors believe represent a continuum, with different entities characterized by distinct embryopathologic features. The indications and surgical management options are discussed for these individual intracranial pathologies with relevant illustrations, and a novel classification is proposed for persistent falcine sinus (PFS). The role of neurointervention and/or microsurgery in specific cases such as sinus pericranii and enlarged emissary veins of the skull is highlighted.

A better understanding of the pathophysiology and developmental anatomy of these lesions can reduce treatment morbidity and mortality. Some patients, including those with vein of Galen malformations (VOGMs), can present with the added systemic morbidity of a high-output cardiac failure. Although VOGM is the most studied and classified of the above-mentioned eDVAs, the authors believe that grouping the former with the other venous anomalies/abnormalities listed above would enable the clinician to convey the exact morphophysiological configuration of these lesions, predict their natural history with respect to evolving venous hypertension or stroke, and extrapolate invaluable insights from VOGM treatment to the treatment of other eDVAs. In recent years, many of these symptomatic venous malformations have been treated with endovascular interventions, although these techniques are still being refined. The authors highlight the broad concept of eDVAs and hope that this work will serve as a basis for future studies investigating the role of evolving focal venous hypertension/global intracranial hypertension and possibilities of fetal surgical intervention in these cases.

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KEYWORDS dural venous sinuses; sinus pericranii; persistent embryonic sinus; vein of Galen malformations; enlarged emissary veins; venous hypertension; embryology; developmental venous anomaly

Many publications over the past few decades have identified the natural history and clinical significance of classic developmental venous anomalies (DVAs) of the brain that are located intraaxially. However, extraaxial DVAs (eDVAs) are discussed in relatively fewer publications, and their various subdivisions are often considered as separate entities, not as part of a spectrum. Classic developmental cranial venous anomalies drain the normal brain parenchyma and are located intraaxially. We believe that persistent embryonic sinuses, sinus pericranii, venous varices or aneurysmal malformations, and enlarged emissary veins comprise a spectrum...
Among the persistent embryonic intracranial sinuses, the most common and hence widely studied is the persistent falcine sinus (PFS). Manoj et al. described how in the early stages of fetal development, when the embryo is around 20 mm in length, the primitive falx cerebri has been assumed to contain anastomotic venous loops, termed the sagittal plexus. As the embryo develops, the primitive falx cerebri will enclose the sagittal plexus. Tubbs et al. hypothesized that the sagittal plexus gives rise to the falcine venous plexus. The dominating venous channel within the sagittal plexus gives rise to the dorsally located superior sagittal sinus, while the inferior sagittal sinus and straight sinus develop from the ventral portion of the sagittal plexus. The smaller channels of the sagittal plexus gradually disappear. When the occipital pole grows posteriorly, caudal anastomotic loops to the superior sagittal sinus and straight sinus allow for development in the same direction. Bartels et al. stated that the falcine sinus and the straight sinus arise from the mesenchyme found within the menencephalic flexure. Sener emphasized that dysfunction in the mesenchyme may contribute to a PFS and/or absent straight sinus. Multiple studies have shown that congenital absence of the straight sinus may cause a predilection for recanalization of the falcine sinus to compensate for an atretic sinus.

The existing anatomical classifications of PFS do not account for associated developmental anomalies of the venous system in the brain; there is no definitive radiological criterion for grading PFS that encompasses the architecture and associations of PFS. Ryu stated that it is important that clinicians be cognizant of the congenital anomalies linked to falcine sinuses or straight sinus occlusion. In the event that a straight sinus occlusion or absence is apparent, it is important to also look for a persisting falcine sinus with a concurrent vein of Galen malformation.
TABLE 1. Manjila grading of persistent falcine sinuses

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description</th>
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<tbody>
<tr>
<td>1</td>
<td>Normal falx w/ PFS disconnected from SSS w/ or w/o focal duplication of SSS</td>
</tr>
<tr>
<td>2</td>
<td>Hypoplastic falx cerebri posterior to PFS w/ or w/o hypoplasia of distal SSS</td>
</tr>
<tr>
<td>3</td>
<td>PFS w/ normal falx; deficient straight sinus w/ or w/o dysplastic tentorium cerebelli</td>
</tr>
<tr>
<td>4</td>
<td>PFS w/ hypoplastic falx/SSS associated w/ deficient straight sinus w/ or w/o dysplastic tentorium cerebelli</td>
</tr>
<tr>
<td>5</td>
<td>PFS grades 1–4 w/ additional neurovascular developmental lesions, like vein of Galen pathologies &amp; enlarged parietal emissary veins</td>
</tr>
<tr>
<td></td>
<td>Subtype A w/ atretic parietal/occipital cephalocele</td>
</tr>
<tr>
<td></td>
<td>Subtype B w/o atretic parietal/occipital cephalocele</td>
</tr>
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</table>

PFS = persistent falcine sinus; SSS = superior sagittal sinus. 
Tenting or peaking of the tentorium can occur with any of the grades. Similarly, an enlarged parietal emissary foramen or a focal duplication of the SSS can appear with or without an atretic parietal/occipital cephalocele.

This prompted us to propose a novel grading system for PFS which covered coexisting lesions in a logical stepwise manner (Table 1).

The falcine sinus, a normal venous structure in the fetus located between the dural leaves of the falx cerebri, is an ascending midline vein that connects the vein of Galen and sagittal sinus above the level of the confluence of sinuses and disappears before birth. Bartels et al. emphasized that the venous plexus in the developing tentorium will influence development of the straight sinus in addition to the sagittal plexus. A PFS with no additional associated pathology is highly unlikely. Rhoton also stated that there is communication between the inferior sagittal sinus and the superior sagittal sinus by way of a venous channel in the falx cerebri. Tubbs et al. validated this and classified the falcine venous sinus based on its relationship with the superior sagittal sinus. Type I falcine sinuses have no communication with the superior sagittal sinus and extensive network of inferior sagittal sinus tributaries; type II falcine sinuses have some degree of communication with the superior sagittal sinus; and type III falcine sinuses have considerable communication with the superior sagittal sinus. These classifications were based on the behavior of the embryonic venous plexus within the falx cerebri, not accounting for adjacent brain changes (such as atretic cephalocele) or associated defects in the straight sinus (Fig. 2).

The newly proposed Manjila classification takes into consideration the pattern of PFS along with associated defects in brain and other dural venous sinuses (straight sinus, superior sagittal sinus); this classification can be used to understand the natural history better and stratify patterns of PFS lesions in syndromic conditions, as they occur with other AVMs, agenesis of the corpus callosum, osteogenesis imperfecta, acrocephalosyndactyly (Apert syndrome), occipital encephaloceles, absent or abnormal tentorium cerebrelli, enlarged parietal foramina, and Chiari II malformations.

Cephaloceles, also known as “occult,” “rudimentary,” or “abortive” cephaloceles, are skin-covered cranial dysraphisms caused by “abnormal disjunction of the mesencephalon,” as proposed by McLone and De Leon. Several groups of authors have stated that cephaloceles are associated with venous anomalies, such as vertical positioning of the embryonic straight sinus, absence of the straight sinus, focal duplication of the superior sagittal sinus, and elongation of the vein of Galen. Brunelle et al. suggested that venous anomalies are seen when the cephalocele is located above the confluence of sinuses. Occipital cephaloceles are the most common type of cephalocele and typically occur between the lambdoid suture and foramen magnum. Bartels and colleagues' study of patients with an occipital encephalocele found an absent straight sinus in all patients as seen on MR venography (MRV). They also observed an alternative drainage pathway of the galenic system through a falcine sinus. Occipital cephaloceles have a much better clinical prognosis than parietal cephaloceles. We feel that this is due to involvement of less eloquent dysmorphic brain and fewer draining veins to the posterior third of the sagittal sinus than to the middle third of the sagittal sinus. Reddy et al. reported a link between prominent parietal foramina and PFS.

Indications and Surgical Management Options

From a surgical standpoint, if the straight sinus is compromised during surgery, the falcine sinus may have a significant role in alternative venous drainage. This crucial information can help in the management of venous hypertension in selected cases—for example, where a PFS is implicated in pseudotumor cerebri. Conversely, an iatrogenic falcine injury can cause major bleeding and should not be missed on preoperative CTA. As Tubbs et al. pointed out, there is a rather safe avascular corridor in the falx cerebri for performing a contralateral transfalcine approach to mesial frontoparietal or medial occipital lesions. They observed that most falcine veins are located in the posterior third of the falx cerebri, especially in the inferior two-thirds of the falx in rostrocaudal dimensions. Caution must be exercised in resection of large falcine meningiomas and techniques required to turn a flap of falx to access the contralateral cerebral hemisphere.

For surgical treatment of a dural arteriovenous fistula (DAVF), resection of the falx cerebri, including the falcine sinus as well as the abnormal falcine plexus, is an appropriate strategy, according to Yamaguchi et al.
cal treatment of atretic cephalocele may be considered if there is reasonable risk for traumatic injury, seizures, and/or bleeding as well as for cosmetic reasons. In many endoscopic cases, such as interhemispheric approaches, inadvertent parafalcine subdural hematomas are found, likely due to falx plexus injury.

Sinus Pericranii

Sinus pericranii (SP) is a rare extradural venous anomaly in which there is an abnormal connection between the extracranial venous system and the intracranial dural sinuses via a connecting diploic vein (Fig. 3). Although the condition is usually asymptomatic, there have also been reports of symptomatic SP causing debilitating headaches, ataxia, nausea, vomiting, hearing loss, epilepsy, bradypnea, and bradycardia. Furthermore, SP can cause fatal complications, such as thrombosis, air embolism after trauma, or massive hemorrhage. SP has also been found to be associated with cases of esophageal atresia, macrocephaly, mental retardation, cerebrofacial arteriovenous malformation type II (CAMS II), meningocoele, craniosynostosis, and other intracranial venous anomalies. Although SP is rare, due to the potential severity of the complications, symptoms, and associated syndromes, understanding the etiology, anatomy, presentation, diagnosis, and treatment yields significant value.

Etiopathogenesis

The comprehension of the possible mechanisms in the formation of SP alerts the clinician as to when to suspect this condition. Several etiologies have been proposed for the SP, some postulated many decades ago, even prior to modern radiological imaging. The best accepted is Mastin’s classification, which categorizes the pathogenesis into 3 etiologies: congenital, spontaneous, and traumatic. Congenital SP can be histologically identified by the presence of vascular endothelium. This supports Muller’s view that congenital hemangiomas are true angiommas coexisting with other vascular anomalies. The fact that there have been cases of SP in which the patients presented with systemic angiommas, persistent trigeminal arteries, cerebrofacial arteriovenous malformation type III (CAMS III), facial hemangioma, solitary DVA, vein of Galen aneurysmal malformation, dural sinus malformation, vein of Galen hypoplasia, and intraosseous AVMs supports this congenital theory. SP associated with such other congenital, developmental, or genetic conditions can be explained by errors in the late stages of embryogenesis because the development of brain parenchyma and arterial vasculature precedes vein development. Congenital SPs have been found in cases of craniosynostosis, such as Apert syndrome, Crouzon syndrome, oxycephaly, and trigonocephaly.
niosynostosis and SP has not been agreed upon, it can be reasoned that increased intracranial pressure plays a role in affecting venous sinus pressure. Renier and Marchac indicated that intracranial hypertension was observed in 42% of complex cases of craniosynostosis.59 Furthermore, in the absence of overt congenital or traumatic origins, venous hypertension has been suggested as an independent etiology for SP.65 This view is supported by the fact that there have been multiple cases of macrocephaly and hydrocephalus coexisting with SP.19,25

Both traumatic and spontaneously acquired lesions of SP have a connective tissue endothelial lining, which differentiates them from congenital SP.15 Therefore, it can be reasoned that spontaneous SPs are most likely the result of minor trauma that went largely unrecognized. Traumatic SP has been caused by outer table fractures of the cranium, venous sinus tears with epidural hematomas or depressed bone fragments, or disruption of an emissary vein as it leaves the skull.65

Clinical Presentation

The vast majority of SPs present as a nontender compressible and fluctuant subcutaneous mass in the midline of the cranium that increases with Valsalva maneuver. However, in the case of thrombosis, the mass will not be compressible.19 Ota et al. reported that SPs are located in the frontal (40%), parietal (34%), occipital (18%), and temporal regions (4%).51 Most cases of SP communicate with the superior sagittal sinus.44

Although an overwhelming majority of SPs present in the midline of the cranium, cases of trauma and some cases of craniosynostosis can be exceptions. Traumatic SP usually occurs in the area of the trauma. The case report of Sadler et al. presents a traumatic SP in the retromastoid area that communicates with the transverse sinus.64 Furthermore, there have been cases of craniosynostosis and coexisting SPs that have occurred in the temporal regions and, in some cases, communicated with the transverse sinus.43 However, Mitsukawa et al. reported 7 cases of craniosynostosis with associated midline SP.43

Diagnosis and Management

The diagnosis of SP can be made based on clinical examination and imaging showing an extracranial sinus communicating with an intracranial sinus. Useful investigations include ultrasonography, plain radiography, CT, MRI, and digital subtraction angiography. Ultrasonography will show a venous vessel crossing a hyperechoic cranial vault.19 Plain skull radiographs and CT scans will show bony defects, cortical thinning, and focal bony erosions.19 CT with intravenous contrast will show an enhanced subcutaneous scalp mass corresponding to the SP in relation to the dural sinus. MRI will also show the flow intensity of the SP and the relationship with the adjacent
intracranial sinus. Digital subtraction angiography can be used to assess the flow properties of the SP and the adjacent intracranial dural venous sinus. In a suspected or diagnosed SP, lack of enhancement, or a filling defect, in any intravenous contrast study suggests thrombosis of the SP. Carpenter et al. found that CT venography was imperative to distinguish intraluminal thrombi from adjacent slow-flowing blood. We feel that a CT venogram study is better to identify an intraluminal filling defect than MR venogram with contrast administration. The radiological differential diagnosis of subepicranial hygromas, traumatic leptomeningeal cysts, epidermoid/dermoid tumors, encephalocele, and venous cavernoma of the scalp should be also entertained in these cases.

If treatment is considered, definitive vascular imaging using venography should be performed to rule out vascular malformations that mimic SP or any DVAs that coexist with SP, and to assess the intracranial venous dynamics. Asano et al. used delayed 3D CTA to differentiate a subepicranial varix, which lacks intracranial communication with a sinus, from an SP. Gandolfo et al. proposed that SP is a cutaneous sign of an underlying DVA. Thus, in consideration of surgery, additional investigation of the intracranial outlet, such as a jugular bulb. Subsequently, the team used an endovascular transvenous approach to embolize the diploic vein of the SP with N-butyl cyanoacrylate glue mixed with 1:1 Lipiodol. Kesler et al. obtained a venogram via direct puncture of the SP followed by microcatheter insertion of two Guglielmi detachable coils and 0.5 ml of acrylic glue injection mixed 4:1 with Lipiodol. Rangel-Castilla et al. used DynaCT to assess the morphological structure of the SP and digital subtraction angiography to evaluate the venous flow patterns. Subsequently, endovascular embolization with hydrocoils and ethylene vinyl alcohol copolymer (Onyx, ev3 Inc./Covidien) was used to achieve fast embolization of the diploic vein and minimize the number of coils used. With recent advances in endovascular therapy, more SP cases are being treated with nonoperative interventions safely and effectively for indications such as pain relief and cosmesis.

Enlarged Emissary Veins

Emissary veins are valveless bidirectional veins that connect the extraaxial venous system with the intracranial sinuses of the brain. They serve essential neurophysiological functions such as draining blood from the scalp, equalizing intracranial pressure during cerebral congestion, temperature regulation, and dominant sinus drainage in cases of venous sinus occlusion or vascular malformation. Emissary veins have also been implicated in major pathophysiological processes. Emissary veins can provide conduits for infection in the intracranial sinuses and brain parenchyma. Enlarged emissary veins have been involved as sources of intraoperative air emboli and have also been described as forming part of an AVF and providing vascular connection to the diploic venous system in the formation of SP. Along with the developmental, pathophysiological, and neurosurgical implications of emissary veins, a detailed understanding of emissary veins is necessary in the context of venous flow patterns and even inheritance patterns. The inheritance pattern of enlarged parietal foramina is autosomal dominant, raising possibilities of inheritance in eDVAs.

Emissary Vein Anatomy and Neurosurgical Implications

The size and location of emissary veins may vary between individuals, and not all emissary veins are found in every person. We used PubMed to identify the most clinically relevant sites of emissary veins. These include, but are not limited to, the condylid emissary vein, mastoid emissary vein, occipital emissary vein, parietal emissary vein, pterosquamosal sinus, ophthalmic veins, sphenoidal emissary vein, emissary veins of the foramen ovale, internal carotid venous plexus, emissary vein of the foramen cecum, emissary vein of the foramen lacerum, emissary

Microsurgical and Endovascular Approaches

In the surgical approach to SP, the goal is to resect the exophytic scalp mass and ligate the emissary vein communicating with the intracranial sinus. Cranietomy in the region of the skull with the involved diploic veins followed by cranioplasty has been used. The removal of the extracranial mass and closure of the transosseous channel using bone wax, absorbable gelatin packing, coagulation, and diamond-drilled bony dust has also been reported in the literature. However, some studies have reported significant hemorrhage with these surgical methods.

A minimally invasive endovascular approach to treatment has been used after angiographic evaluation of the intracranial venous drainage patterns. Brook et al. used angiography to demonstrate independent drainage of a DVA into the superior sagittal sinus without connection to the SP. Subsequently, the team used an endovascular transvenous approach to embolize the diploic vein of the SP with N-butyl cyanoacrylate glue mixed with 1:1 Lipiodol. Kesler et al. obtained a venogram via direct puncture of the SP followed by microcatheter insertion of two Guglielmi detachable coils and 0.5 ml of acrylic glue injection mixed 4:1 with Lipiodol. Rangel-Castilla et al. used DynaCT to assess the morphological structure of the SP and digital subtraction angiography to evaluate the venous flow patterns. Subsequently, endovascular embolization with hydrocoils and ethylene vinyl alcohol copolymer (Onyx, ev3 Inc./Covidien) was used to achieve fast embolization of the diploic vein and minimize the number of coils used. With recent advances in endovascular therapy, more SP cases are being treated with nonoperative interventions safely and effectively for indications such as pain relief and cosmesis.

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Development and Anomalies of Emissary Veins

The early embryonic brain includes superficial, middle, and deep capillary venous plexus layers. The superficial vessels drain into the external jugular vein and the middle and deep layers drain into the internal jugular vein. Emissary veins will have residual connections with the superficial and middle layers by the third trimester. The emissary veins begin most of their development after ballooning of the transverse sinus. The emissary veins of the hypoglossal canal are the first to develop. The earliest emissary veins will drain medially into the primitive dural venous sinuses. The last emissaries to develop are the parietal emissary veins. The chondrocranium will develop around these vessels and will form respective foramina. As the jugular sinuses grow, most emissary veins disappear, but some persist and enlarge. Chauhan et al. proposed that this may be caused by a failure of normal maturation of the sigmoid-jugular sinuses.

Enlarged emissary veins can form from compensatory physiological mechanisms and pathophysiological mechanisms. The absence of valves in emissary veins can result in enlargement during times of increased intracranial pressure. It has also been proposed that emissary veins regulate the temperature of the brain and dilate in hyperthermia and constrict in hypothermia. Additionally, an enlarged emissary foramen does not always transmit a large emissary vein (Fig. 4).

Pathological enlargement of emissary veins can result from aberrant connections to arteries or venous varicces, and enlarged emissary veins have been associated with AVFs and extraaxial venous malformations. Dilated mastoid emissary veins, mastoid canals, anterior condylar emissary veins, and hypoglossal canals have been found with AVFs. In regard to the pathogenesis of DAVFs and enlarged emissary veins, Miyachi et al. proposed that local inflammation near an emissary vein’s penetration of the cranium results in enlargement, neovascularization, and subsequent aberrant connections to nearby arteries. In a case of SP, there has been enlargement of the parietal emissary vein, which served as the connection to the diploic venous system. Emissary veins can also enlarge secondary to destruction of primary venous outlets in craniofacial syndromes.

Emissary veins also have a wide range of neurosurgical implications. They can serve as landmarks to underlying venous sinuses. Enlarged emissary veins can be used in endovascular procedures to access intracranial sinuses, and can cause hemorrhage with avulsion from the bone in trauma and surgery; also, improper coagulation can cause extensive thrombosis of intracranial sinuses. In lateral approaches to the cranial base, sphenoidal emissary veins and veins of the foramen ovale should be appreciated. The connection of the cavernous sinus with the petrosquamous sinus may result in complications in epidural procedures due to its connections with the foramen ovale emissary veins and Vesalius emissary veins. In petrosectomy operations, the location of the petrosquamosal emissary vein should be identified and avoided. Postoperative venous epidural hematoma has been reported to occur from bleeding emissary veins. Careful consideration should be used in the coagulation of bleeding emissary veins. Temporalis muscle plugs have been shown to be efficient coagulation agents, while bone wax has the risk of propagating into deeper intracranial sinuses and causing thrombosis and even pulmonary embolism.

Emissary veins can also provide essential conduits in complex endovascular neurosurgical procedures. The mastoid emissary vein has been used to gain endovascular access to the transverse sigmoid sinus for the treatment of DAVFs. In a similar endovascular approach, Schipper et al. and many others have used the supraorbital vein to gain access to the ophthalmic vein and subsequently to the cavernous sinus. Collateral emissary veins have been shown to take the main venous outflow in the removal of cranial base tumors. Preoperative angiography including a delayed venous phase is necessary to anticipate and avoid a catastrophic thrombosis and postoperative brain infarction.

FIG. 4. Enlarged parietal foramen in a 2-week-old female infant. Axial T2-weighted images show a focal defect in the right parietal bone with some extrusion of the meningeal covering and prominence of the CSF space. Of note, no abnormal vascular communication is present.
or as a result of increased collateral flow secondary to tumor burden. In an anecdotal report, an enlarged condylar emissary vein was found during the removal of a hypoglossal neurinoma. The coagulation of this channel resulted in a dural venous sinus thrombosis and subsequent cerebellar infarction. Therefore, it is recommended that a preoperative angiographic evaluation be performed to avoid these surgical complications. Emissary veins can be primarily evaluated on MRI, CT, cerebral angiography, and ultrasonography; any of these modalities could be used for serial follow-up studies, while more invasive catheter-based angiography may be reserved for periprocedural evaluations.

### Venous Varices and Aneurysmal Malformations

Venous varices can occur in the brain as a result of dural sinus occlusion or as a part of AVMs or DAVFs. There is an etiologic role of venous hypertension and sinus thrombosis in the former and cortical venous reflux in the latter. However, an intraaxial DVA with a coexistent varix would suggest a causative role for venous hypertension during embryogenesis and changes in posterior projection of the developing brain. The prototypical case for complex extraaxial venous varix remains the vein of Galen malformation (VOGM), often called vein of Galen aneurysmal malformation (VGAM).

### Vein of Galen Aneurysmal Malformations

The vein of Galen is a venous structure within the quadrigeminal cistern formed by the confluence of the internal cerebral veins and the basal veins of Rosenthal. A vein of Galen malformation (VOGM) is a rare vascular anomaly caused by abnormal development of the median prosencphalic vein of Markowski, which is a precursor of the vein of Galen, and VOGMs are thought to develop by the 11th week of gestation. Raybaud et al. have made significant contributions to the embryological understanding of the vein of Galen. VOGMs, due to the lack of a nidus, are sometimes referred to as arteriovenous fistulas (AVFs),
with the first vein of Galen arteriovenous malformations (VGAMs) being reported by Steinheil. The term VGAM will be used for the remainder of this discussion.

VGAMs typically occur in young children and can be diagnosed in utero by ultrasound and/or fetal MRI. With respect to the chronological development of VGAMs, there is debate as to the timing; they have been described as arising between the 6th and 8th to 11th weeks of gestation. Venous drainage from VGAMs is variable, as there are a number of potential outflow pathways, which is also dependent on the venous anatomy that is or is not present at that stage of development. Examples include a normal straight sinus, a hypoplastic straight sinus, an absent straight sinus, and/or a persistent falcine sinus (PFS). The shunting associated with VGAMs is associated with high-output cardiac failure during the neonatal period. Positive outcomes are most likely to occur if endovascular therapy is initiated in a timely manner, as Lasjaunias has emphasized, and a delay may also cause disruption in cerebrospinal fluid flow.

Yaşargil, Litvak et al., and Lasjaunias and colleagues established various VGAM classifications as shown in Table 2 and Fig. 5. Litvak et al. described the first classification system in 1960, but it lost popularity once newer classification systems were introduced. The newer VGAM classification systems—the Lasjaunias classification—and the Yaşargil classification—are broadly based on the origin (and number) of the feeding arteries and the location of the arteriovenous communication and are useful due to their prognostic and/or treatment implications. Endovascular embolization is the current standard of care for the treatment of VGAMs. While transvenous coil embolization, glue embolization, and the combination of the two are becoming popular treatment options, transarterial embolization is often preferred. The goal of endovascular therapy is to occlude the vessels that supply the arteriovenous shunt or to embolize vessels that drain the lesion itself. The treatment strategy and outcome depend primarily on the venous anatomy, the classification of the lesion, and the presence or absence of cardiac failure (Fig. 6).

We would like to point out that eDVAs that present as cisternal venous varices do not have a natural history similar to that of venous angiomas or intraaxial DVAs. In contrast to venous angiomas (associated with intraaxial DVAs), which are dilated veins of developmental origin, the intraaxial DVA complexes can cause obstructive hydrocephalus in certain cases. Venous angiomas are rarely symptomatic in intraaxial locations, although they can cause hemorrhages and infarctions in some of these cases. Nonetheless, there are no indications for surgical interventions in venous angiomas. Hydrocephalus resulting from an eDVA is seldom reported.
Conclusions

We have provided a contemporary update on the various extraaxial developmental cranial venous anomalies with relevant classifications and clinical implications. Persistent embryonic sinuses, sinus pericranii, venous varices or aneurysmal malformations, and enlarged emissary veins, considered as a continuum, form the spectrum of extraaxial DVAs (eDVAs). Symptomatic eDVAs are being treated with endovascular interventions alone or in combination with microsurgery, although these techniques are still being refined. Basic knowledge of developmental anatomy and natural history of these lesions offers an ideal research substrate for future studies looking at the role of evolving venous hypertension/intracranial hypertension and possibilities of fetal surgical intervention in these cases.

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Disclosures
The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

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