INTRODUCTION

Vein of Galen aneurysmal malformation (VGM) is a rare congenital abnormality with a prevalence of less than 1/25,000 deliveries. It is characterized by a midline intracranial vascular fistula with aneurysmal dilatation of the vein of Galen. During the normal development, the large median prosencephalic vein of Markowski gradually involutes and eventually persists as the great cerebral vein of Galen. Thus, VGM is assumed to be a persistent structure of the embryonic median prosencephalic vein of Markowski.

Majority of the VGM present in the early childhood, and the most striking presentation is congestive cardiac failure in the newborn period. In the past, this was usually associated with rapid progression to multisystem organ failure and death, despite vigorous treatment and supportive medical care. With the advent of endovascular techniques, treatment outcome of VGM has much improved. Embolization of feeding arteries or draining veins can considerably reduce the shunt flow. The authors were to present their experience of treatment for VGM, mainly in terms of endovascular embolization.

MATERIALS AND METHODS

Overall, six patients were diagnosed and treated in our institute between January 1998 and December 2010. We retrospectively reviewed medical record and imaging data of them. VGMs were angiographically classified according to the Yasargil’s method (type I, small pure cisternal fistula between VGM and pericallosal arteries or posterior cerebral arteries; type II, multiple fistulous communications between the vein of Galen and the thalamoperforating vessels; type III, high flow mixed type I and II). VGMs were treated with observation and endovascular embolization via the transvenous and transarterial approaches, using various detachable coils (Guglielmi detachable coils [Boston Scientific, Fremont, CA, USA], MicroPlex [MicroVention, Aliso Viejo, CA, USA], Trufill-DCS [Cordis, Miami Lakes, FL, USA], Axium [ev3, Irvine, CA, USA]), fibered pushable coils (Tornado, Cook, Bloomington, IL, USA), 25-33% N-butyl...
2-cyanoacrylate glue (NBCA, B Braun, Melsungen, Germany), and ethylene vinyl alcohol copolymer dissolved in dimethyl sulfoxide (Onyx®, ev3, Irvine, CA, USA). Transvenous approach was performed via the femoral and internal jugular veins to VGMs. Transarterial approach was done via the femoral artery toward the various arterial feeders such as anterior and posterior choroidal (AChA and PChA) and pericallosal arteries (PA). Clinical outcome was graded according to Jones’ 5-point scale (score of 0 : death; score of 1 : severe neurological impairment; score of 2 : moderate neurological impairment; score of 3 : mild neurological impairment; score of 4 : normal)³).

RESULTS

Clinical and angiographic data were summarized in Table 1. All patients were boys under two years old with mean age of 4.4±5.7 months (range, 0.2 to 15 months) at the time of the initial treatment. Two patients (patient 1 and 2) were delivered by cesarean section in normal full term, three (patient 3 to 5) were by vaginal delivery in normal full-term, and one (patient 6) were by vaginal delivery on gestational age of 35 weeks. Clinical presentation included asymptomatic cardiomegaly, cardiac anomalies such as arterial septal defect (ASD), patent ductus arteriosus (PDA), and patent foramen ovale (PFO), congestive heart failure, increased head circumference (HC), and hydrocephalus. Three asymptomatic patients were detected on brain ultrasound shortly after birth. His HC was 95 percentile and cardiac auscultation revealed a VGM. Subsequent cerebral angiography showed that the lesion had multiple direct fistulae from lenticulostriate artery, AChA, and PChA to the vein of Galen. VGM was embolized twice with multiple coils via transvenous approach. First embolization was performed without any complication. However, patient died after the second embolization due to subarachnoid and intraventricular hemorrhages (Fig. 1).

Illustrative cases

Patient 1

A six-month child presented with cold sweating and feeding difficulty. Brain ultrasonography and magnetic resonance imaging revealed a VGM. Subsequent cerebral angiography showed that the lesion had multiple direct fistulae from lenticulostriate artery, AChA, and PChA to the vein of Galen. VGM was embolized twice with multiple coils via transvenous approach. First embolization was performed without any complication. However, patient died after the second embolization due to subarachnoid and intraventricular hemorrhages (Fig. 1).

Patient 6

A two-week child was born by vaginal delivery on gestational age of 35 weeks with 2.9 kg. There were no perinatal problems, and VGM was coincidentally found on screening brain ultrasonography shortly after birth. His HC was 95 percentile and cardiomegaly was identified on echocardiography. It was supplied

Table 1. Clinical data and treatment outcomes in patients with vein of Galen malformation

| No  | Sex/Age (month) | Clinical presentations | Type* | Arterial feeders | Treatment (frequency) | Embolization materials | Complication | Treatment result | Follow-up duration (year) | Clinical outcome |
|-----|----------------|-----------------------|-------|------------------|----------------------|-----------------------|--------------|-----------------|---------------------------|-----------------|
| 1   | M/6            | CHF, HCP              | III   | AChA, PChA, LSA  | TVE (2)              | Coils                 | SAH+IVH      | Incomplete      | 0                         | 0               |
| 2   | M/4            | CHF, HC†, ASD         | III   | PA, PChA         | TVE (7)              | Coils                 | NBCA, SAH+IVH| Incomplete      | 0                         | 0               |
| 3   | M/1            | CHF                   | III   | PChA, PCoA, PA, TPA | TAE (5)            | Coils, NBCA, Onyx†   | None         | Incomplete      | 6                         | 3               |
| 4   | M/15           | PFO                   | II    | PChA, TPA        | Observation          | None                  | None         | Spontaneous regression | 1                       | 4               |
| 5   | M/0.4          | CM, PDA               | II    | PChA             | TAE (1)              | Coils                 | NBCA         | Incomplete      | 3                         | 4               |
| 6   | M/0.2          | HC†, CM               | III   | PChA, AChA, PA, TPA | TAE (2)            | Coils, Onyx†         | None         | Complete        | 2                         | 4               |

*Yasargil’classification of vein of Galen malformation⁴. Clinical outcome was scored according to an article⁵: CHF : congestive heart failure, HCP : hydrocephalus, AChA : anterior choroidal artery, PChA : posterior choroidal artery, LSA : lenticulostriate artery, TVE : transvenous embolization, SAH : subarachnoid hemorrhage, IVH : intraventricular hemorrhage, HC : head circumference, ASD : arterial septal defect, PA : pericallosal artery, NBCA : N-butyl 2-cyanoacrylate glue, PCoA : posterior communicating artery, TPA : thalamoperforating artery, TAE : transarterial embolization, PFO : patent foramen ovale, CM : cardiomegaly, PDA : patent ductus arteriosus
by PChA, AChA, PA, and TPA. Transarterial embolization via PChA was incompletely performed at two weeks of age with coils with no complication. The second transarterial embolization via PChA achieved complete occlusion of the lesion at one year of age with Onyx with no complication (Fig. 2). Thereafter, he has been followed-up without any problems.

**DISCUSSION**

VGMs consist of high-flow arteriovenous fistulas and can lead to congestive heart failure and hydrocephalus. They are rare, and usually associated with high mortality and morbidity, particularly in neonates (up to 100%). There have been a lot of efforts to understand the angioarchitecture and pathophysiology, and to perform effective treatment. As the endovascular techniques have developed, understanding of the VGM has greatly advanced during the past two decades. Analyzing the vascular anatomy of VGM and correlating it with embryologic development of the cerebral vasculature, Raybaud et al. concluded that the malformation develops between the sixth and eleventh week of gestation, after development of the circle of Willis. It is thought to result from the development of an arteriovenous connection between primitive choroidal vessels and the median prosencephalic vein of Markowski.

Clinical presentations and angioarchitectures are known to have a tight correlation with age at presentation. Symptomatic neonates typically present with severe cardiac failure, and high flow shunt of Yasargil type I is dominant. Infants usually present with increase in head circumference, hydrocephalus, or seizure. Older children tend to present with milder symptoms such as headache. In the present study, age did not have such a close relationship with the clinical presentations. Two (patient 1 and 2) of three patients with congestive heart failure (CHF) were older than one month of age, and one (patient 3) of three neonates presented with CHF. It was because the age at admission did not exactly correspond to the one at clinical presentations. Diagnosis of VGM in patient 2 was delayed because initial symptoms had been thought to result from ASD alone. Patient 5 and 6 were incidentally diagnosed with routine check-ups of brain ultrasonography.

Endovascular intervention is known to be treatment of choice for VGM. There are two kinds of techniques: transvenous and transarterial embolization. Transvenous approach is performed, either with femoral or jugular vein, or through direct puncture of the torcular. The venous approach to the VGMs is technically less demanding and can be performed in situations where a superselective arterial embolization is not feasible. However, the drawback of transvenous approach is that rapid closure of the venous drainage can cause the development of "normal perfusion breakthrough syndrome", with severe complications, including malignant brain swelling and intracranial hemorrhages. In addition, recent studies reported that normal deep veins could be connected to the venous component of a VGM. This particular anatomy can be a major pitfall of transvenous embolization because it may potentially result in detrimental complications such as venous infarct and hemorrhage. There-
fore, majority of VGMs have been treated with transarterial embolization except just a few cases. Casasco et al. achieved excellent results with transvenous approach and believed that transvenous approach was better than transarterial embolization or surgery. They could escape from the disastrous complications in the manner that embolization stopped when intra-VGM pressure increased more than 50% of the initial pressure. In the present study, patients who were treated with transvenous approach died all due to the deep brain swelling and intracranial hemorrhage. The reasons of selecting a transvenous approach were that they had complex angioarchitectures of Yasargil type III. Thereafter, transarterial approach with or without combined transvenous approach was the treatment of choice for the treatment of VGMs in our institute.

On the other hand, transarterial embolization is performed through the perforators. Some fine perforators are sometimes impossible to approach, and embolic material within a larger fistula can fly away from the fistula point and occlude the draining veins, which can cause venous hypertension. As Onyx has been used, transarterial embolization became more feasible and safer. It is a bio-compatible liquid polymer that precipitates and solidifies upon contact with blood, thus forming a soft, spongy embolus. So, it can be molded and extended through the fine channels distant from the injection point for a longer time rather than the previous glues. In this series, transarterial embolization showed satisfactory clinical outcome even after incomplete treatment. Furthermore, procedures with Onyx were safe and effective that one case could achieve complete occlusion. As the clinical experience of Onyx increases, better treatment results are expected.

This study has a critical limitation. Case number is too small to draw an analytic conclusion but an individual description. Although VGM itself is extremely rare, the true incidence may be higher because VGMs are known to consist of 30% of all pediatric vascular anomalies. Some factors may contribute the rarity of the disease, such as low birth rate, unreported treatment at the other institutes, preterm detection and abortion, and undetected cases. In spite of the rarity, the intention of this report was to share the experience of disastrous situations with treatment strategies for VGMs.

CONCLUSION

VGMs are rare, but serious diseases with high morbidity and mortality. As the endovascular procedures are introduced, treatment of VGMs is successfully performed. In this series, transarterial embolization showed better outcome than transvenous approach. However, small number of case and ongoing advancement in endovascular techniques require more experiences and research.

References
1. Casasco A, Lylyk P, Hodes JE, Kohan G, Aymard A, Merland JJ: Percutaneous transvenous catheterization and embolization of vein of galen aneurysms. Neurosurgery 28 : 260-266, 1991
2. Gaillard P, O’riordan DP, Burger I, Lehmann CU: Confirmation of communication between deep venous drainage and the vein of galen after treatment of a vein of Galen aneurysmal malformation in an infant presenting with severe pulmonary hypertension. AJNR Am J Neuroradiol 27 : 317-320, 2006
3. Gaillard P, O’riordan DP, Burger I, Levrier O, Jallo G, Tamargo RJ, et al.: Diagnosis and management of vein of galen aneurysmal malformations. J Perinatol 25 : 542-551, 2005
4. García-Monaco R, De Victor D, Mann C, Hannedouche A, Terbrugge K, Lasjaunias P: Congestive cardiac manifestations from cerebrocranial arteriovenous shunts. Endovascular management in 30 children. Childs Nerv Syst 7 : 48-52, 1991
5. Jones BV, Ball WS, Tomsick TA, Millard J, Crane KR: Vein of Galen aneurysmal malformation: diagnosis and treatment of 13 children with extended clinical follow-up. AJNR Am J Neuroradiol 23 : 1717-1724, 2002
6. Khullar D, Andejeani AM, Bulsara KR: Evolution of treatment options for vein of Galen malformations. J Neurosurg Pediatrics 6 : 444-451, 2010
7. Lasjaunias PL, Chng SM, Sachet M, Alvarez H, Rodesch G, Garcia-Monaco R: The management of vein of Galen aneurysmal malformations. Neurosurgery 59 : S184-S194; discussion S3-S13, 2006
8. Lasjaunias P, Hui F, Zerah M, Garcia-Monaco R, Malherbe V, Rodesch G, et al.: Cerebral arteriovenous malformations in children. Management of 179 consecutive cases and review of the literature. Childs Nerv Syst 11 : 66-79; discussion 79, 1995
9. Long DM, Seljeskog EL, Chou SN, French LA: Giant arteriovenous malformations of infancy and childhood. J Neurosurg 40 : 304-312, 1974
10. Park SH, Hwang SK, Kim YT, Kim SL: Vein of Galen malformation in a neonate: case report. J Korean Neurosurg Soc 34 : 51-53, 2003
11. Raybaud CA, Strother CM, Hald JK: Aneurysms of the vein of Galen: embryonic considerations and anatomical features relating to the pathogenesis of the malformation. Neuroradiology 31 : 109-128, 1989
12. Yamashita Y, Abe T, Ohara N, Maruoka T, Toyoda O, Inoue O, et al.: Successful treatment of neonatal aneurysmal dilatation of the vein of Galen: the role of prenatal diagnosis and trans-arterial embolization. Neuroradiology 34 : 457-459, 1992
13. Yasargil MG: Microneurosurgery IIIIB. New York : Thieme Medical Publishers, 1988, pp 323-357