Cerebellar Glioependymal Cyst

El Kim

Department of Neurosurgery, Dongsan Hospital, Keimyung University School of Medicine, Daegu, Korea

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Correspondence
El Kim
Department of Neurosurgery, Dongsan Hospital, Keimyung University School of Medicine, 1035 Daegubeol-daero, Dalseo-gu, Daegu 42601, Korea
Tel: +82-53-528-7823
Fax: +82-53-528-7356
E-mail: bach1158@dsmc.or.kr

INTRODUCTION

Glioependymal cysts (GECs) are benign neuroepithelial lesions that have been infrequently detected; they are also known as neuroglial cysts. These developmental lesions virtually occur everywhere within the brain and spinal cord, and represent less than 1% of primary intracranial cysts [1]. They might virtually present at intraparenchymal, intraventricular, subarachnoid, intraneural, and intraspinal regions [2-4]. The reported intracerebral GECs are predominantly originated in the central white matter of the frontal lobe [5]. For this reason, there is a lack of available information about GECs of the infratentorial compartment in the relevant literature [6,7].

The author herein illustrates an interesting case harboring a GEC developed in the cerebellum near the fourth ventricle. The present report also provides the histogenesis, radiographic findings, and pathological details for differentiating this rare type from other cysts more common in the posterior cranial fossa.

CASE REPORT

This 36-year-old female with cystic lesion in the right cerebellum was hospitalized for evaluating headache and dizziness. Brain images showed a well-defined, ovoid mass adjacent to the fourth ventricle. After drainage and excision of the cyst, the patient became symptom free. Pathology examination disclosed low cuboidal epithelium and glial cells in the cyst wall. The radiological features, neurological manifestations, and the operations for GECs of the present localization are described in this short communication.

Key Words Glioependymal cyst; Neuroepithelial cyst; Cerebellum.
The posterior cranial fossa is an extremely rare site for GECs that lead to clinical manifestations. To the best of the author's knowledge, only 11 cases with these cysts pathologically diagnosed have been referenced in the related literature (Table 1) [8-14]. Six cysts were observed in the cerebellopontine angle region, three at the midbrain, and two entirely within cerebellar hemisphere. No cysts involved the cavity of the fourth ventricle. The author confirmed that most cases were women and their average age was 42.3 years. And, those of the prior reports are cysts with various sizes seen in adult presentation.

The cysts are congenital lesions, arising from rests of embryonic neural tube segments that become sequestered within the developing white matter [2,15-17]. These ectopic cells thus evolve into epithelial-lined cysts that are located intracerebrally, extracerebrally or anywhere along the neural axis and do not communicate with the ventricular system. Consequently, almost all cases reported have exhibited a propensity for the periventricular location within the supra- and infra-tentorial compartments for their given cysts [14].

They usually contain cerebrospinal fluid-like clear fluid; how-
ever different aspects of intracavitary contents (xanthochromic, opalescent, milky, and turbid) have been described in the reports [1,18]. In pathological evaluation, they reveal the partly ciliated cuboidal or columnar lining epitheliums resting directly over the astroglial tissue bundles without an intervening basal membrane [19]. The immunohistochemistry depicts that the surface-lining cells are positively stained for glial fibrillar acidic protein and S-100 protein, and are negative for epithelial membrane antigen, cytokeratin, and carcinoembryonic antigen [8,16]. This specificity suggests that GECs have an origination from the neural ectoderm. Both ependymal cyst and choroid plexus cyst have the similar characteristics to the current cyst in the immunohistochemistry, so that it is important to notion that the expression of the glial stroma is a decisive to accurately find out the GEC from other nontumorous cysts.

Mostly, GECs are clinically silent; however, in certain cases, they cause symptoms and require treatment. Due to the secretory nature of the epithelial cells, these cysts can grow to sizable masses over time and progressively result in the signs and symptoms [20]. Generally, the neurologic deficits depend on the locations and expansion of the GECs. The adult supratentorial cysts have presented with a varying symptomatology [21-23]. The posterior fossa lesions more commonly have induced dizziness, double vision, syncope, gait disturbance, cranial neuropathy, and hemifacial spasm [6-9]. Characteristically, those presentations were linked to the grade of obstruction of the cerebrospinal fluid flow, severity of intracranial hypertension, and degree of compression of the neural-vascular structures including the brain stem, cerebral aqueduct, cerebellum, and the fourth ventricle.

Typically, CT scans depicted unilocular cyst with thin-walled morphology [5]. On MR images, the cysts have smooth border and their contents do not restrict in diffusion weighted sequence [7,24]. In radiological differentiation from GECs, scalloping of the adjacent cranium is often seen in the arachnoid cysts [25]. However, the most difficult lesion to distinguish from the GEC is an arachnoid cyst, especially when it occupies the cerebellopontine angle. The contents of epidermoid and dermoid cysts were positive with diffusion weighted MRI and heterogeneous on fluid attenuated inversion recovery image. Furthermore, typically both grow larger with insinuating pattern [26]. Next, porencephalic cysts communicate with the lateral ventricle and show surrounding gliosis and spongiosis [27]. Contrary to GECs, enlarged perivascular spaces are typically multiple and cluster around the basal ganglia. Infectious cyst, such as neurocysticercosis, is commonly smaller than 10 mm and partly calcified and enhances [19]. Neurenteric cysts are characterized for isointensity to hyperintensity on T1-weighted MRI [6]. Lastly, ependymal cyst and choroid plexus cyst have the similar characteristics to the current cyst in the immunohistochemistry, so that it is important to notion that the expression of the glial stroma is a decisive to accurately find out the GEC from other nontumorous cysts.

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The operative approaches are selected according to the cyst locations and their proximity to subarachnoid or ventricular spaces. Numerous procedures have been introduced, including trephination and aspiration, craniotomy and excision, fenestration into the cistern, open or endoscopic cystoventriculostomy, and cystoperitoneal, cystocisternal or cystoventricular

![Table 1. Reported adult cases of glioependymal cysts in the posterior fossa](image)

| Study | Age/sex | Location | Cyst size | Symptomatology | Therapy | Follow-up | Outcome |
|-------|---------|----------|-----------|----------------|---------|-----------|---------|
| Ho and Chason [8] | 47/F | CPA cistern | 30 mm | Dizziness | Total excision | 37 months | Symptom free |
| Tsuchida et al. [9] | 29/M | Cerebellum | 27×31×23 mm | Dizziness | Total excision | 38 months | Symptom free |
| Harada et al. [10] | 27/F | CPA cistern | 30 mm | HFS | CSF and MVD | 10 months | Symptom free |
| Shenouda et al. [11] | 39/M | CPA cistern | Cryptic | HFS | Excision and MVD | Unknown | Symptom free |
| Endo et al. [12] | 54/F | Midbrain tegmentum | 27×22 mm | Headache, blurred vision, HDC | ECV and ETV | 3 years | Symptom free |
| Park et al. [13] | 44/F | CPA cistern | 66×54 mm | Syncope | Excision and CSF | 4 months | Symptom free |
| Prieto et al. [14] | 30/M | Midbrain tectum | 18×28 mm | Gaze palsy, headache, HDC | ECV and VPS | 6 months | Symptom free |
| This report | 36/F | Cerebellum | 25×23×22 mm | Headache, dizziness | Total excision | 15 years | Symptom free |

CPA, cerebellopontine angle; HFS, hemifacial spasm; HDC, hydrocephalus; CSF, cystosubarachnoid fenestration; MVD, microvascular decompression; ECV, endoscopic cystoventriculostomy; VPS, ventriculoperitoneal shunt; ETV, endoscopic third ventriculostomy.
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shunting [1,13,22]. Recently, endoscopic fenestration of the cerebral hemispheric cysts thorough the adjacent ventricle is done as the first line of treatment for the selected cases to avoid the craniotomy and shunt dependence [12,23,28]. Simple puncture and drainage by using the open or stereotaxic has a higher recollection rate of the cysts [29]. In light of such evidences, complete resection is the best mode of treatment whenever possible, especially for symptomatic cerebellar GECs at an early stage of the disease. Interestingly, six reports have described GECs of the cerebellopontine angle cistern, three of which presented with hemispheric spasms and they underwent the total or partial excision with microvascular decompression, resulting in long-term resolution of symptoms [10,11]. For the cysts in the dorsal midbrain with hydrocephalus, adding third ventriculostomy, aqueductal stenting or ventriculoperitoneal shunt might to lessen the possible risk for recurrence in cerebrospinal fluid circulation disturbance.

In summary, the presented reporting highlights that the cerebellum is an unusual location for intraparenchymal GECs. Despite of its rarity, this congenital condition should be considered a potential diagnosis for the cysts either midline or at cerebellopontine angle in the posterior fossa. Complete cystectomy is curative operation for cerebellar GECs with neurological symptoms and sings.

Conflicts of Interest

The author has no potential conflicts of interest.

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None

ORCID ID

El Kim https://orcid.org/0000-0002-7664-6030

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