Pediatric Giant Cell Glioblastoma Presenting with Intracranial Dissemination at Diagnosis: A Case Report

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Abstract

Giant cell glioblastoma (GCG) is a rare subtype of glioblastoma multiforme (GBM), and it often occurs in younger patients; however, its onset in children is extremely noticeable. A 7-year-old girl presented with a headache and restlessness. A giant tumor that was 7 cm in diameter was found by magnetic resonance imaging (MRI) in the left frontal lobe with intracranial dissemination. Because the tumor had extended to the lateral ventricles and occluded the foramen of Monro causing hydrocephalus, she underwent ventricular drainage and neuro-endoscopic biopsy from the left posterior horn of the lateral ventricle. The initial pathological diagnosis was an atypical teratoid/rhabdoid tumor (AT/RT). When the dissemination subsided after the first chemotherapy with vincristine, doxorubicin, and cyclophosphamide, she underwent the first tumor resection via a left frontal transcortical approach. After surgery, the second chemotherapy with ifosfamide, cisplatin, and etoposide was not effective for the residual tumor and intracranial dissemination. The second surgery via a transcallosal approach achieved nearly total resection leading to an improvement of the hydrocephalus. The definitive pathological diagnosis was GCG. Despite chemoradiation therapy, the dissemination in the basal cistern reappeared and the hydrocephalus worsened. She was obliged to receive a ventriculo-peritoneal (VP) shunt and palliative care at home; however, her poor condition prevented her discharge. Ten months after admission, she died of tumor progression. The peritoneal dissemination was demonstrated by cytology of ascites. In conclusion, although unusual, pediatric GCG may be disseminated at diagnosis, in which case both tumor and hydrocephalus control need to be considered.

Keywords: giant cell glioblastoma, pediatrics, dissemination, ventriculo-peritoneal shunt, immunohistochemistry

Introduction

Malignant gliomas such as glioblastoma multiforme (GBM) are relatively rare in children compared to adults.1 Malignant gliomas are generally considered more common in children; however, only 0.6%–7.9% of all GBMs occur during childhood.3 More than 70% of pediatric GBMs occur in the second decade; thus, pediatric GBMs in the first decade are especially unusual.4 Giant cell glioblastoma (GCG) is an unusual GBM subtype defined by the latest classification of the World Health Organization.5,6 GCG constitutes only 1% of adult GBMs and 3% of pediatric GBMs.6,7 Although GCGs are diagnosed at relatively younger ages than the more common GBM,7 they rarely develop in childhood.

We present a case of pediatric GCG with intracranial dissemination, which is extremely rare with respect to age and presence of dissemination at
diagnosis. We discuss the radiological diagnosis, treatment, and pathological examination of this disease.

Case Report

History and preoperative examination

A 7-year-old Japanese girl presented with complaints of headaches since the age of 3 years. One month prior to admission, she had posterior neck pain and loss of appetite; subsequently, she visited a nearby hospital. A cervical magnetic resonance imaging (MRI) examination did not reveal any significant changes (Fig. 1A). However, 1 week later, she complained of a headache and loss of appetite again. She then had a head MRI examination, which showed a left frontal mass that was 7 cm in diameter with dissemination into the left lateral ventricular wall and the basal cistern (Figs. 1B–1D).

She did not have any previous developmental disorders or other medical history. When she was admitted to our hospital, the patient's Glasgow Coma Scale score was 10 (Eyes [E]=2, Verbal [V]=4, Motor [M]=4), but she had no apparent motor weakness. Gadolinium-enhanced head MRI revealed the heterogeneously enhanced tumor in the left frontal lobe extending to the lateral ventricles with acute hydrocephalus due to obstruction of the foramen of Monro. At first, the patient underwent ventricular drainage and a neuro-endoscopic biopsy via the left posterior horn of the lateral ventricle. The initial pathological diagnosis was an atypical teratoid/rhabdoid tumor (AT/RT). Fortunately, the first 14-day chemotherapy cycle with vincristine (1.5 mg/m²), doxorubicin (37.5 mg/m²), and cyclophosphamide (1800 mg/m²) (VDC) reduced the intracranial dissemination. Next, she underwent the first tumor resection via a left frontal transcortical approach. Afterward, she underwent a second 21-day chemotherapy cycle with ifosfamide (2000 mg/m²), carboplatin (500 mg/m²), and etoposide (100 mg/m²) (ICE); however, the residual tumor showed no changes. Instead, the hydrocephalus worsened, and the dissemination was newly detected in the spinal region (Fig. 2) and cerebellopontine angle. Approximately 1.5 months after the first resection, she underwent a second tumor resection via a transcallosal approach, and a nearly total resection was achieved, resulting in improvement of the hydrocephalus. Then, the patient received radiation therapy: 23.4 Gy for the craniocerebral lesion, 30 Gy for the tumor cavity, and 29.6 Gy for the cerebellopontine lesion. Four cycles of weekly vincristine therapies (1.5 mg/m²) and a second VDC therapy were not effective. After the patient was diagnosed with GCG by the Central Pathological Diagnosis of the Department of Pathology, National Center for Child Health and Development, she received chemotherapy with five cycles of temozolomide (150 mg/m²) and four cycles of bevacizumab (10 mg/kg), which unfortunately did not help to control the dissemination. As the hydrocephalus worsened, she was obliged to undergo a ventriculoperitoneal (VP) shunt instead of ventricular drainage for subsequent postoperative palliative home care. Eventually, she could not be discharged home because of anuresis and oral feeding difficulties. Two months after the VP shunt, she experienced severe abdominal distension, followed by dyspnea. Ten months after admission, she died of respiratory failure caused by tumor progression. The peritoneal dissemination was demonstrated by cytology of ascites.

Histological examination

Hematoxylin–eosin staining of the first surgical specimens showed giant cells consisting mainly of a mega nucleus and a bizarre polynucleus, which was sometimes located eccentrically, as well as cells with a wide range of eosinophilic, vacuolated, and

![Fig. 1 (A) Plain cervical MRI scan performed at the previous hospital showing no abnormal findings. (B–D) Initial contrast-enhanced head MRI scan showing a homogeneously enhanced 7-cm left frontal mass lesion (A), and dissemination into the left lateral ventricular wall (B) and the basal cistern (C). MRI: magnetic resonance imaging.](image)
foamy cytoplasm (Fig. 3A). Tumors consisted of small to medium spindle-shaped or multi-ridged cells, cells with multipolar processes, and small cells with a high nuclear-to-cytoplasm ratio (Fig. 3B). These cells were diffusely and densely proliferating. Many small- to medium-sized cells undergoing mitosis were observed, but only a few giant cells undergoing mitosis were observed. Hypervascularization of small vessels with congestion was observed, but no microvascular proliferation was detected. In some areas, geographic necroses were observed.

Immunohistochemistry showed diffuse positive nuclear staining for p53 (1:200, Dako, Fig. 3C) and INI-1 (1:500, Becton Dickinson, Fig. 3D), and cytoplasmatic staining for vimentin (1:200, Leica, Fig. 3E). No immunostaining was detected for IDH1R132H (1:100, Dianova, Fig. 3F), H3K27M (1:200, Abcam, Fig. 3G), and glial fibrillary acidic protein (GFAP) (1:100, Dako, Fig. 3H). MIB-1 immunostaining (1:200; Dako, Fig. 3I) was positive in 30% of giant cells and 60% of small cells. In addition, using the Pyrosequencing method from the Japan Children’s Cancer Group, we obtained the results that both H3F3A K27M and H3F3A G34 were wildtype. When taken together, this case was IDH-wildtype, p53 mutant, and H3F3A K27M and H3F3A G34 wildtype. These findings were considered to correspond to a diagnosis of GCG. Immunohistochemistry for PTEN and CD133 was performed to investigate factors related to intracranial dissemination in the early stage. As a result, PTEN (1:10000; proteintech, Fig. 3J) was strongly positive in the cytoplasm of the tumor cells; therefore, PTEN mutation was denied. In contrast, moderate CD133 expression (1:1500; proteintech, Fig. 3K) was observed in not only giant cells but also small cells. Papanicolaou staining for the ascites collected by peritoneal puncture revealed a lot of small oval atypical cells rather than giant cells (Fig. 3L).

Discussion

GCG is included in the World Health Organization classification as a rare subtype of GBM. The mean age of onset for GCG is lower than that for GBM, and GCG commonly localizes to the frontal and temporal lobes. The average age at diagnosis is 59 years for GBM and 44 years for GCG. Therefore, the present case represents a very young case in the GCG group. The H3K27M mutation is frequent in diffuse intrinsic pontine glioma and non-brain stem midline tumor, while G34R/V mutation occurs in pediatric high-grade glioma (HGG) of the cerebral cortex. It was reported that the former occurred mainly in younger patients (median age 11 years), whereas the latter occurred in older patients (median age 20 years). Accordingly, in terms of age and site of origin, there is no contradiction that both mutations were wildtype in this case. Based on molecular diagnosis, p53
Fig. 3  (A) Hematoxylin–eosin staining of the first surgical specimens showing giant cells mainly consisting of a mega nucleus and a bizarre polynucleus, which was sometimes located eccentrically, and cells with a wide range of eosinophilic, vacuolated, and foamy cytoplasm. (B) Hematoxylin–eosin staining showing clusters of small oval cells. (C–K) Immunohistochemical staining for the first surgical specimens. (C) Diffuse positivity in the nuclei for p53. (D) Diffuse positivity in the nuclei for INI1. (E) Diffuse positivity in the cytoplasm for vimentin. (F) Negative staining for IDH1R132H. (G) Negative staining for H3K27M. (H) Negative staining for GFAP. (I) Positive staining for MIB-1 in 30% of giant cells. (J) Positive staining for PTEN in the cytoplasm of giant cells. (K) Positive staining for CD133 in both giant (arrowheads) and small cells (arrows), contrast negative findings in endothelial cells and inflammatory cells. (L) Papanicolaou staining for the ascites collected by peritoneal puncture at the time of death showing some atypical oval cells with intense hematoxylin staining (arrows); these cells resemble small cells shown in B and K. Scale bar indicates 100 µm. GFAP: glial fibrillary acidic protein.
The MIB-1 LI of up to 60% in our case might have been a factor associated with dissemination. Altogether, the factors considered to be related to intracranial dissemination in this case were GFAP-negative, p53 mutation, CD 133-positive, and high MIB1-LI.

Korshunov et al.\(^{40}\) reported that >90% of cases displaying tumor dissemination through the central nervous system at recurrence were found in the high-risk group, which supports the very poor prognosis of the present case of GCG with initially presenting dissemination. In contrast, Benesch et al.\(^ {31}\) reported that the median overall survival was not significantly different between pediatric patients with and without dissemination of HGG. However, because the prognosis cannot be improved only by removing the tumor mass lesion, a treatment strategy for dissemination is needed from an early stage of therapy. Accordingly, we performed a first round of chemotherapy (VDC therapy) before the first craniotomy, followed by a second round (ICE therapy) and a second craniotomy. The chemotherapies with VDC and ICE were chosen based on the initial diagnosis of AT/RT. The former showed a partial response for dissemination while the latter was ineffective. Gross total resection was achieved by the two-staged surgery, and the final pathological diagnosis was changed to GCG. When we changed the chemotherapy treatment to temozolomide, we had difficulty in controlling the dissemination. As a result, the hydrocephalus worsened, and the patient could not continue with the ventricular drainage. Then, we performed a VP shunt so that the patient could receive palliative care at home. We repeatedly examined the cytology of the cerebrospinal fluid collected from the ventricular drainage before the VP shunt surgery, because of a previously reported brain tumor seeding following shunt placement.\(^ {41}\) Unfortunately, a small number of atypical cells were detected in the cytology immediately before the shunt placement, but surgery was performed because of the patient’s limited life expectancy. As a result, the headache subsided; however, the patient’s level of consciousness worsened, and abdominal distension occurred 1 month after the shunt placement. The cytology of ascites did not reveal any giant cells, but some small oval atypical cells, which resembled the cells observed by hematoxylin–eosin staining in the first surgical specimens. Hoffmann et al.\(^ {42}\) used a Millipore filter encased in Rickham reservoir for VP shunt to avoid pleural metastasis. The use of such a filter is likely to cause obstruction of the shunt tube, but may be necessary to prevent pleural metastasis, since a VP shunt may be a potential route for iatrogenic metastasis.\(^ {43}\)
Conclusion

We describe an extremely rare case of GCG as it developed at a very early age, where the patient presented with intracranial dissemination at the time of diagnosis. Even though GCG has a better outcome than GBM, the prognosis for GCG with dissemination is generally poor. When VP shunt placement is performed to treat hydrocephalus, the possibility of pleural metastasis should be monitored closely, and the timing of the surgery could play a role in palliative care.

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Conflicts of Interest Disclosure

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