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Syed Sarmad Bukhari
_Aga Khan University_, sarmad.bukhari@aku.edu

Muhammad Junaid
_Bahria University Medical and Dental College, Karachi, Pakistan_

Ali Afzal
_Bahria University Medical and Dental College, Karachi, Pakistan_

Anisa Kulsoom
_Fauji Foundation Hospital, Jhelum Road, Rawalpindi, Punjab, Pakistan_

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Case Report

Primary pediatric cerebellar gliosarcoma

Syed Sarmad Bukhari1, Muhammad Junaid2, Ali Afzal2, Anisa Kulsoom3

1Department of Neurosurgery, Aga Khan University Hospital, 2Department of Neurosurgery, Bahria University Medical and Dental College, Karachi, 3Department of Radiology, Fauji Foundation Hospital, Jhelum Road, Rawalpindi, Punjab, Pakistan.

E-mail: *Syed Sarmad Bukhari - sarmadbukhari@gmail.com; Muhammad Junaid - junaidkhan1034@gmail.com; Ali Afzal - dokterali@gmail.com; Anisa Kulsoom - docanisa@hotmail.com

ABSTRACT

Background: Primary gliosarcomas of the central nervous are rare and very few have been reported in the infratentorial compartment. Here, we describe such a lesion in a 12-year-old male.

Case Description: A 12-year-old male presented with headache, ataxia, and vomiting. When Magnetic resonance studies documented a posterior fossa lesion, he underwent placement of a right ventriculoperitoneal shunt followed by a suboccipital craniectomy. The lesion proved to be a primary gliosarcoma. Unfortunately, it recurred 2 years later and required repeated resection.

Conclusion: Here, we reviewed the rare case of a 12-year-old male requiring shunt placement and suboccipital craniectomy for a primary gliosarcoma that recurred 2 years later.

Keywords: Gliosarcoma, Pediatric, Primary

INTRODUCTION

Gliosarcomas are malignant brain tumors that usually involve the temporal lobe in the supratentorial compartment; only rarely do they involve the cerebellum. They typically contain mixed glial and mesenchymal elements. The present WHO system classifies them as a variant of glioblastomas. Here, we present a 12-year-old male who first underwent ventriculoperitoneal shunt placement and a suboccipital craniectomy for a primary cerebellar gliosarcoma resection; unfortunately, the tumor recurred 2 years later.

CASE DESCRIPTION

Clinical and magnetic resonance (MR) presentation

A 12-year-old male presented with headache, ataxia, and vomiting of 3 months duration. The brain MRI with/without contrast revealed a heterogenous appearing lesion within the fourth ventricle measuring 3.18 × 3.26 × 3.90 cm. There was also a small cystic component noted along the right lateral aspect which was isointense on T1, iso to hyperintense on T2, and hyperintense on the FLAIR study. Anteriorly, the tumor was causing marked compression of the medulla oblongata, pons, and midbrain; this resulted in obstructive hydrocephalus.
Surgery

The patient first underwent placement of a right-sided ventriculoperitoneal shunt. This was followed by a suboccipital craniotomy for subtotal excision of tumor. At surgery, the lesion was firm pseudo-encapsulated, and partially adherent to the brainstem on the left; due to the adhesions, a portion of the left-sided lesion was unresectable.

Pathology and immunohistochemistry

The microscopic features and the immunohistochemical profile revealed a heterogeneous tumor with hypercellular tissue that contained spindle-shaped cells that were vimentin positive. These findings confirmed the diagnosis of a cerebellar gliosarcoma [Figure 1]. The patient was, therefore, referred to oncology for further management.

Postoperative course and repeat MR 1 year later

The patient did well post-surgery for 1 year. However, when the MRI was repeated, it documented a significant recurrent lesion now measuring 4.1 × 3.3 × 3.7 cm [Figures 2 and 3]. However, there was no increase in hydrocephalus due to the presence of the VP shunt. In addition, the MR showed herniation of the cerebellum tonsils through the foramen magnum with attendant syringohydromyelia involving the cervicodorsal cord. Although the patient's family was offered further surgery, they declined, and the patient was lost to follow-up.

Poor outcome 2 years later without further surgery

The patient next seen 2 years postoperatively. Over the prior 3 months, he had developed progressive quadriplegia and was wheelchair bound for 1 month. Although a follow-up MRI was requested, only a T2 sequence could be obtained due to financial constraints [Figure 4]. Due to very significant brainstem compression, the patient underwent a repeated suboccipital resection of tumor. Despite the subsequent postoperative MRI showing maximum safe resection of tumor, the patient's clinical condition did not improve in the early postoperative period (1 month of follow-up) [Figure 5].

DISCUSSION

WHO classification

Gliosarcoma has been classified by the World Health Organization (WHO) as high-grade tumors (Grade IV) resembling glioblastomas. They account for nearly 2% of Grade IV glial neoplasms and have a similar age distribution as glioblastoma.\[3\]

Location

Most, if not all, cases of primary gliosarcomas in the literature occur in the supratentorial compartment, very rarely, they involve the cerebellum in adults, and even less frequently, in children.\[3,7\] These tumors typically involve the peripheral regions of the temporal lobe followed by the frontal and parietal lobes; they only occasionally involve the corpus callosum. Here, we reported the unusual presentation of a cerebellar gliosarcoma occurring in a 12-year-old male within the fourth ventricle, abutting the brainstem.

Computed tomography (CT) and MR findings of gliosarcoma

CT

CT findings are similar to those for an infiltrating glioblastoma. They may appear as a well-defined hyperdense mass with heterogeneous ring enhancement due to their fibrous component. Central necrosis is less common due to their predilection for peripheral lobe locations. Dural involvement is not uncommon.\[5,6\]

MR

Nitta et al.\[6\] reported that the MRI appearance of infratentorial gliosarcomas resembles those of supratentorial gliosarcomas, for example, peripheral location within the

Figure 1: (a-c) Photomicrographs with H&E staining demonstrating the cytoarchitecture of the tumor with hypercellular and hypocellular myxoid areas. The cellular areas are composed of spindle shaped cells having mildly pleomorphic elongated nuclei. Scattered eosinophilic globules with few microtubules are seen as well.
cerebellum, accompanied by a broad meningeal base, with multifocality. Further, on magnetic resonance imaging, gliosarcomas appear as a heterogeneous mass both on T1- and T2-weighted images, with strong peripheral enhancement, and central hemorrhage or necrosis.[5]

Pathology of gliosarcomas

Pathologically, gliosarcomas are characterized by a biphasic tissue pattern with alternating areas of glial and mesenchymal differentiation.[4] The sarcomatous component shows malignant spindle cells with features of fibrosarcoma or malignant fibrous histiocytoma. Areas of mesenchymal differentiation into cartilage, bone, fat, and smooth or skeletal muscle may be an associated finding. In our case, the tumor was predominantly sarcomatous and had a mixture of clearly malignant mesenchymal GFAP-negative areas and GFAP-positive glial areas. Furthermore, collagen deposition and reticulin fibers were seen in the sarcomatous component which helped differentiate it from glioblastoma (e.g., with florid fibroblastic proliferation).[2]

Local recurrence

Local recurrence and metastases of gliosarcomas along cerebrospinal fluid pathways are common.[8] Of interest, extracranial metastases carry an even worse prognosis.[6] Common differential diagnoses for gliosarcomas include glioblastoma, metastasis, brain abscess, or other infratentorial intraxial tumors (e.g., hemangioblastomas or medulloblastomas).[3,5]
CONCLUSION

Primary infratentorial gliosarcomas are rare in children and are particularly located within the infratentorial compartment. Although shunting and resection may result in short-term improvement, these lesions typically recur, resulting in devastating neurological deficits and short-term survival.

Declaration of patient consent

Patient's consent not required as patients identity is not disclosed or compromised.

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Conflicts of interest

There are no conflicts of interest.

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