INTRODUCTION: Cerebellar glioblastoma (cGBM) is extremely rare, accounting for 0.7–0.9% of all gliomas. Few studies have reported on clinical course, histopathology, and prognosis. In this report, we discussed cases which were diagnosed as cGBM, and were treated in our institute. Materials and Methods. We retrospectively analyzed 9 cGBMs (age ranged 41 to 85 years, median 69), operated at our institute after 2010 January, and evaluated their <MGMT> promoter methylation, <IDH1> mutation, and Copy Number Variation status detected by methylation-specific PCR (MSP), DNA sequencing or immunohistochemistry, and Multiplex Ligation-dependent Probe Amplification (MLPA), respectively. RESULTS: All patients underwent resection; 3 gross total resections (GTRs, 33%), 2 subtotal resections, 4 partial resections, with relatively low achievement of GTR. The tumor location predominated in the cerebellar hemisphere (7 patients, 78%) over vermis (2). One patient had brain stem invasion. After surgery, 8 patients received temozolomide (TMZ) and radiotherapy (RT), while only one did RT alone. After recurrence, three patients were treated with bevacizumab monotherapy, and other three received either TMZ and RT, TMZ and ACNU, or TMZ monotherapy. The median progression-free survival (PFS) was 12.0 months, and the median overall survival (OS) was 17.1 months. Five patients (<MGMT> methylated, whereas all were <IDH1>-wild type, <PTEN> deletion was negative in all patients). <EGFR> amplification and combination <PDGER> amplification and <CDKN2A> deletion were found in one patient each. DISCUSSION: Despite the lower rate of GTR, there was a tendency of longer PFS compared to supratentorial GBM (cGBM). The clinical course after recurrence was unfavorable, and OS was similar to that of cGBM. cGBMs appeared to lack the typical genetic mutations occurred in GBM, suggesting that cGBMs might be stimulated with different regulatory cellular signals.

CS-01 GIAN T CELL TUMOR IN THE SKULL BASE BONE TREATED WITH ANTI-RANKL INHIBITOR

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Gi ant cell tumor of bone is a rare and osteolytic neoplasm that usually affects the epiphyses in long bones of the extremities. They seldom occur in the skull, preferentially affecting the sphenoid and temporal bones. Most pathologically benign, and total removal by surgery was regarded as the first treatment, however, it was very difficult in skull lesion. In 2014 the molecular targeting drug anti-RANKL inhibitor was approved in Japan. We report a case in which an anti-RANKL inhibitor was administered to a skull base bone giant cell tumor that was difficult to remove completely. A 56-year-old man with a sudden right neck pain followed by dysphoria and dysphagia was referred to our hospital. Computed tomography showed 4.4 x 2.0 cm osteolytic lesion involving the right occipital bone and occipital condyle. Magnetic resonance imaging demonstrated an extensive soft-tissue mass occupying. Surgical biopsy was performed and the pathological diagnosis was giant cell tumor. Patient received the anti-RANKL inhibitor (Denosumab)®. After 4 weeks, resolution was observed, and neurological symptoms were improved after 12 weeks. Patient has been on good course for 5 years without recurrence and is still following-up.

CS-02 CLINICAL AND MOLECULAR ANALYSIS OF ASTROBLASTOMAS

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Astroblastoma is extremely rare brain tumor which mostly arise in cerebral hemisphere of children and young adult. Limited data exists on its clinical feature and molecular analysis. We recently experienced two female patients with astroblastoma in the cerebrum.

Case 1 is a 3-year-old girl. She developed left hemiparesis. CT and MRI revealed large supratentorial mass with cystic component and calcification. Gross total removal was achieved. She is well without recurrence on MRI one year after surgery. Case 2 is 42-year-old lady. She developed partial seizure. CT and MRI revealed a mass with ring-enhancement in the left temporal lobe. Gross total removal was achieved under awake craniotomy. She is well without recurrence on MRI six months after surgery. Pathologic examination of both patients showed pseudorosette formation of tumor cells around vasculature. Molecular analysis revealed rearrangement of MN-1 in case 1 but not in case 2. Case 2 showed BRAF V600E mutation and loss of CDKN2A/2B. Both patients received no adjuvant therapy.

Prognosis of astroblastoma varies and standard of treatment is not established. Gross total resection is associated with increased survival, but the role of adjuvant chemotherapy and radiation therapy are controversial. Advances in molecular analysis will lead to establish molecular classification and risk-adapted treatment strategy.

CS-03 LARGE CYSTIC INTRADURAL SCHWANNOMA IN CERVICAL REGION: A CASE REPORT

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Schwannomas are the most common intradural extramedullary spinal tumors. However, they are usually solid tumors, and totally cystic changes are rare. Herein, we report a case of a 46-year-old male presenting with numbness of right limbs, right hemiplegia, and posterior neck pain for one year. MRI revealed a well-defined cystic long-segment, from C1 to C6, intradural extramedullary mass. The lesion showed hypointense on T1WI, hyperintense on T2WI, hypointense on DWI, and it was marginally enhanced on the contrast image with Gd-DTPA. C1 laminectomy and hemi-laminectomy from C2 to C6 was performed for tumor resection. The tumor was found to be totally cystic and tended with a jelly-like content. It was completely resected with the attachment of the C3 dorsal root. Histopathological examination confirmed it to be a schwannoma. The mechanism of cyst formation in schwannoma is considered as results of ischemic necrosis associated with tumor growth, or cystous due to degeneration of Antoni type B region. The long-segment, totally cystic intradural cervical schwannoma is rare, but it should be included in the differential diagnosis of a cystic mass in the spinal region. It can be difficult to distinguish cystic spinal schwannomas from other cystic lesions like arachnoid cyst, epidermoid cyst, and neurenteric cyst. Contrast enhanced MRI is useful by enhancing the margin of the tumor.