Corpus Callosal Oligodendroglioma: A rare entity
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Abstract
Introduction: Oligodendroglioma’s (ODG) are primary glial brain tumors that are divided into classical and anaplastic ODG’s. Common pathologies encountered in the corpus callosum include primary central nervous system lymphomas and glioblastoma. Corpus callosal ODG is a rare entity. Case Report: A 54 year old male presented with headache since 6 months and weakness of right side of the body since 4 months. Radiological imaging was suggestive of a heterogeneous minimally enhancing mass arising from the corpus callosum and extending bilaterally into the frontal lobes (left more than right). Left frontal craniotomy and tumor decompression was done. Final histopathological report was “oligodendroglioma grade II but likely to behave in an aggressive manner”. Patient received adjuvant chemo-radiation and on his latest follow up two years post-surgery, radiological imaging showed only gliotic changes in the brain. Conclusion: Corpus callosal oligodendroglioma is a rare entity. The treatment for anaplastic and aggressive ODG’s should include gross total resection (wherever possible) or tumor decompression followed by adjuvant chemo-radiation.

Keywords: Brain Neoplasms, Corpus callosum, Frontal Lobe, Glioma, Oligodendroglioma.
clusters of monomorphic tumor cells, showing classical oligodendrogial like morphology, were seen. The tumor cells classically demonstrated perinuclear halos giving a “fried egg appearance’. Mini-gemistocytes were also seen. The tumor cells were negative for glial fibrillary acidic protein (GFAP) staining. The MIB index was 5-7% [Fig.3]. Significant number of tumor cells showed combined 1p and 19q deletions. The final histopathological report was “oligodendroglioma grade II but likely to behave in an aggressive manner”.

Post-surgery, patient received adjuvant concurrent chemo-radiation followed by 18 cycles of oral temozolomide chemotherapy. On his latest follow up, 2 years post-surgery, CECT and CEMRI scans of the brain showed no obvious enhancing mass lesion but only post treatment gliotic changes in the brain [Fig.1,2].

![Fig.1](image1.png)
**Fig.1:** Pre-operative contrast enhanced computed tomography scan (CECT) of the brain showing a heterogeneous, minimally enhancing, iso to hyper-dense mass lesion involving the corpus callosum and extending into bilateral frontal lobes (left more than right). Few areas of calcification are also noted. Post-operative and post chemo-radiation CECT scan of the brain showing no obvious enhancing mass lesion with only post treatment gliotic changes in the brain.

![Fig.2](image2.png)
**Fig.2:** Pre-operative contrast enhanced magnetic resonance imaging (CEMRI) scan of the brain showing a heterogeneous, minimally enhancing mass lesion involving the corpus callosum. Post-operative and post chemo-radiation CEMRI scan of the brain showing only gliotic changes in the brain.

![Fig.3](image3.png)
**Fig.3(A):** Hematoxylin and eosin stain (4X): showing sheets and papillaroid clusters of monomorphic tumor cells. (B): Hematoxylin and eosin stain (20X): These tumor cells show classical oligodendrogial like morphology with “fried egg appearance”. Mini-gemistocytes are also seen. (C): GFAP staining (20X): Tumor cells are negative for GFAP. (D): MIB staining (20X): MIB index is 5-7%.
Discussion

Oligodendroglioma is believed to originate from the oligodendrocytes of the brain or from a glial precursor cell. These tumors occur in both sexes, with a male: female ratio of 2:1. There are two peaks of incidence: between the age of 6 to 12 years in children and between the age of 35 to 44 years in adults [2]. More than 90% of ODG’s arise in the supra-tentorial white matter, most commonly in the frontal and temporal lobes. Less than 10% occur in the posterior fossa and spinal cord [2]. ODG’s typically demonstrate a slow, infiltrative growth pattern. Cerebrospinal fluid (CSF) metastasis reportedly occurs in up to 14% of the cases [3,4]. In these patients, seizures are the presenting symptom in approximately 50-80% of the cases [3]. Other presenting symptoms include headache, mental status changes, nausea, vomiting, vertigo, visual complaints and/or focal weakness [2]. On diagnostic imaging, ODG’s appear as mass lesions with fairly well defined margins, usually located in the cortex and subcortical white matter [5]. Calcifications are common and are seen in 90% of ODG’s on CT scans [3]. On magnetic resonance spectroscopy (MRS), the lesion demonstrates an increase in choline (Cho) levels with elevated choline/creatine ratio. Positron emission tomography (PET) scan has been reported to be able to differentiate between low grade astrocytoma’s and ODG’s [6]. Derlon et al. in his study found that both these tumor types exhibit glucose hypo-metabolism but on methionine (MET) PET imaging, the uptake of MET was high in the oligo-dendroglioma lesions and decreased, normal or only moderately increased in the astrocytoma group [6]. PET scan may also be useful for grading ODG’s non-invasively [7]. In a different study by Derlon et al. demonstrated that anaplastic ODG’s exhibited a higher 18F-fluro-deoxy-glucose and methionine (MET) uptake than low grade ODG’s [7].

On histological examination, the tumor cells contain uniformly round, homogenous nuclei and a clear cytoplasm. This typical “fried egg appearance” is actually an artifact of formalin fixation which is not present on frozen section and may make the diagnosis of ODG difficult on being frozen [8]. ODG’s may also display a dense network of branching capillaries which is described as “chicken wire” vascular pattern [8]. When the tumor cells invade the grey matter structures such as the cortex, these neoplastic oligodendrocytes tend to cluster around the neurons exhibiting a phenomenon known as “perineuronal satellitosis”. ODG’s usually do not stain for GFAP.

Features associated with anaplastic ODG’s (WHO grade III) and which differentiate these lesions from low grade ODG’s are contrast enhancement on CT or MRI; endothelial proliferation; pleomorphism; tumor proliferation (high mitotic count or high MIB-1 index) and astrocytic component. The high frequency of co-deletion of chromosomal arms 1p and 19q is a striking feature of this glial tumor and is considered as a “genetic signature” of ODG’s. 1p and 19q co-deletion has been associated with both chemosensitivity and improved prognosis in ODG’s [9]. The phosphatase and tensin homolog (PTEN) gene mutations on chromosome 10 are associated with a poor prognosis.

Surgery is the mainstay of treatment for ODG’s. The aim of surgery is gross total resection of the tumor, if possible. The extent of resection largely depends on the location of the tumor and its proximity to ‘eloquent’ areas of the brain. In most studies, the authors have concluded that more complete resection is associated with increased patient survival [10-12]. Post-surgery, radiation therapy is recommended for all patients with a diagnosis of anaplastic ODG regardless of the extent of resection. Radiation therapy is also recommended for patients of grade II ODG who have undergone an incomplete tumor removal or subtotal resection or biopsy of the lesion. Oligodendroglioma’s are chemosensitive tumors.
Post-surgery, adjuvant chemotherapy with either PCV (procarbazine, CCNU or lomustine and vincristine) regimen or oral temozolomide is the standard of care [2]. PCV chemotherapy regimen was the most commonly used chemotherapy regimen to treat ODG’s but is now been superseded by oral temozolomide therapy. This is because of the ease of administration and fewer adverse effects with oral temozolomide therapy.

ODG’s, like all other infiltrating glioma’s have a high rate of recurrence and gradually increase in grade over time. Recurrent tumors are treated with re-surgery (if possible), more aggressive chemotherapy and radiation therapy. Median survival time for grade II ODG’s is 11.6 years and for anaplastic ODG’s is 3.5 years [13]. Factors associated with better prognosis in patients of ODG are age less than 40 years at the time of diagnosis; frontal lobe ODG; gross total resection of the tumor; lower grade of malignancy; good post-operative performance scores; early presentation of the tumor with seizures and combined deletion of chromosomal arms 1p and 19q. Pure ODG’s have a better prognosis than mixed oligoastrocytoma’s.

**Conclusion**

Corpus callosal oligodendroglioma is a rare entity with only a single case reported worldwide till date to our knowledge [1]. The treatment for anaplastic and aggressive ODG’s should include gross total resection (wherever possible) or tumor decompression followed by adjuvant concurrent chemo-radiation. This should be followed by oral temozolomide chemotherapy which in our case has given a good result at the end of 2 years post-surgery.

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