Glioblastoma multiforme subterfuge as acute cerebral hemorrhage: A case report and literature review

Seidu A. Richard,1,3 Yunxia Ye,4 Hao Li,1 Lu Ma,1 Chao You1
1Department of Neurosurgery, Post Graduate Training Centre, West China Hospital, Sichuan University, China; 2Department of Immunology, Jiangsu University, China; 3Department of Surgery, Volta Regional Hospital, Ho, Ghana; 4Department of Pathology, West China Hospital, Sichuan University, China.

Abstract

Hemorrhagic related Glioblastoma multiforme (GBM) are rare and characterizes with severe clinical scuffle. The etiology of this presentation although not well known is believed to be multifactorial. We present a case as well as review on the pathogenesis of evolution of the hematoma into ring enhancing features of GBM on imaging studies. We present a case of 28 years old man who suddenly went into coma for 9 hours preceded with seizures that lasted for 10 minutes. He had no focal neurological signs. CT-Scans images indicated acute cerebral hemorrhage near the frontal horn of the left ventricle with brain edema about the hemorrhagic lesion and MRI done a week later revealed a cerebral ring enhancing lesion. The lesion was partially resected during surgery and immunohistochemical staining confirmed GBM (WHO, grade 4). The diagnosis of intratumoral hemorrhage in GBM was very challenging at the initial stages but with time the hematoma evolved into ring enhancing images typical of GBM. It’s not every intracranial hematoma that is of pure vascular origin.

Introduction

Hemorrhagic related Glioblastoma multiformes (GBMs) are rare and characterizes with severe clinical challenges.1-3 GBM is the most common and lethal type of astrocyte-derived tumor, constituting 50% of adult primary brain tumor cases, followed by anaplastic astrocytoma which constitutes 30% and astrocytoma which also constitutes 20%.4 Licata et al.5 established that 33% of intratumoral bleed is from GBM. The etiology of GBM-related bleeding although not well known; however, it is certainly multifactorial and factors related to this pathogenic process can be categorized into predisposing, initiating, and aggravating.1 This factors can either be local or systemic.1 While the significant local factors comprise of the state of blood vessels and capillaries, the most important systemic factor is coagulopathy which can also occur at the local level and normally maintain the bleeding process.1 The clinical manifestations of cerebral hemorrhage are not exceptionally the first clinical suggestion of a tumor and frequently convey a modest outcome.1,6-8 First hemorrhagic episode in GBM can directly lead to the death of the patient although subsequent bleeding carries a high risk.1 Therefore GBM-related hemorrhage is a very significant risk factor that can result in death of the patient. We present a case of glioblastoma that subterfuge as acute cerebral hemorrhage and review literature on GBM-related hemorrhages with more focus on the pathophysiology as well as pathogenesis of the hemorrhage.

Case Report

We present a case of 28-year-old man who suddenly went into coma for 9 hours preceded with seizures that lasted for 10 minutes. He was initially managed in a periphery hospital before presenting at our facility. The seizures where ionic clonic in nature with no fever, vomiting or micturition. He had no focal neurological signs. Past medical history was unremarkable. Physical examination revealed normal cranial nerves. Power as well as reflexes were present and normal. Routine laboratory as well as other ancillary investigation were normal.

Referral images indicated acute cerebral hemorrhage near the left frontoparietal lobe extending to the frontal horn of the left ventricle with brain edema around the hemorrhagic lesion and midline shift of the brain to the right (Figure 1). The hematoma was mostly centralized at the left parietal lobe. He was admitted in the local hospital for one (1) week prior to referral. MRI done a week later at our facility revealed a cerebral ring enhancing lesion of about 4.6×3.4×4.0 cm in diameter at the left frontoparietal lobe which appear hypo-dense on T1 and hyper-dense on T2. On FLAIR, however the images showed slightly mixed signals with no significant enhancement after administration of contrast agent. There is significant peripheral edema and swelling of the adja-
difficult to different from the normal brain tissue. We excised a chunk of the of lesion and secured total hemostasis. The bone flap was replaced with screws and plates and the subcutaneous tissues, temporalis muscle as well as skin was closed in layers. Postoperative CT-scan revealed total evacuation of the hematoma (Figure 4). Postoperative management was uneventful. Immunohistochemical staining of sample at the pathology department revealed tumor cells that are GAFP (+), Oligo2 (+), EMA (-), IDH-1 (+), ATRX (+), P53 (+) and 40% Ki-67(+) rate. The findings above is consistent with GBM (WHO, grade 4) (Figure 5). The sample was found to be MGMT gene methylation promoter and TERI gene mutation was seen in progress. The patient was discharged home 10 days after surgery and radiotherapy as well as chemotherapeutic sessions initiated a month after surgery. The patient is currently doing very well with no obvious complication.

Discussion

Hemorrhagic related GBMs are rare and characterizes with severe clinical scuffle.1-3 GBM is the most common and lethal type of astrocyte-derived tumor, constituting 50% of adult primary brain tumor cases, followed by anaplastic astrocytoma which constitutes 30% and astrocytoma which also constitutes 20%.4 Licata et al.5 established that 33% of intratumoral bleed is from GBM, 21% from metastases, 13% from anaplastic gliomas, 12% from low-grade gliomas, 12% from meningiomas, 5% from adenomas, 2% from hemangioblastomas, 2% from melanomas, 1% from neuroblastoma and 1% from pinealoblastoma. Kondziolka et al.9,10 observed in their study that out 264 cases with GBM; 17 of these cases constituting about 6.4% had macroscopic hemorrhage while 34 of these cases constituting about 12.9% had microscopic hemorrhage. We also present a case of GBM that subterfuge as acute cerebral hemorrhage. The hemorrhage in our case is a mixed type although predominantly macroscopic hemorrhage.

The etiology of GBM-related bleeding is not well known; however, it is certainly multifactorial and factors related to this pathogenic process can be categorized into predisposing, initiating, and aggravating.1 Atypical new perforating blood vessels that traverse necrotic areas are believed to be the initiation points of intratumoral hemorrhage in GBM.11,12 Also, intratumoral hemorrhage in GBM can occur when large vessels are invaded by the tumor resulting in weakening and breakdown of their walls.11,13

![Figure 1. Referral axial CT-scan images the patient brought from the local hospital. These images show hematoma formation.](image1)

![Figure 2. MRI image done at our facility. A-C) are axial, coronal and sagittal T1 image respectively. D,E) are axial T2 images while 2F is FLAIR. These images show ring like lesions that are consistent with Glioblastoma. The delay window of the evolution process was up to 14 days.](image2)
Furthermore, weak tumor vessels may not have joined the glial meshwork very well hence result in decreased resistance to the trimming forces of the brain.\textsuperscript{11,14} Nonetheless, endothelial proliferation with successive occlusion of the lumen or incidence of intratumoral arteriovenous fistulae are alternative origins of intratumoral hemorrhage.\textsuperscript{11,15} Coagulopathy which occur systemically or at the tumor level usually maintain the bleeding process.\textsuperscript{1} Therefore, making the attainment of hemostasis at hematoma very difficult if conservative treatment options are adapted or during even during surgery.\textsuperscript{11,16}

The clinical manifestations of GBM can differ significantly based on the location and size of the lesion as well as the anatomic structures around the affected brain.\textsuperscript{17-19} Headaches are quite common and unilateral with characteristically progressive in severity.\textsuperscript{20} Considerably increased intracranial pressure may manifest as papilledema if the disease is not noticed early.\textsuperscript{20} Also, larger lesions with substantial mass effect may present with gait imbalance and urinary and/or fecal incontinence. Furthermore, depending on the location of the lesion, focal signs like hemiparesis, sensory loss, or visual field disturbances may occur and rarely, the manifestation of symptoms may be abrupt, masquerading as hemorrhagic stroke.\textsuperscript{17,20} Nevertheless, etymological glitches may be misunderstood for confusion or delirium.\textsuperscript{20} Seizures which may have focal onset occurs in about 20-40% of cases prior to presentation.\textsuperscript{20,21} Therefore, clinically significant hemorrhages related to GBM are comparatively very rare.\textsuperscript{1,22}

Contrast radiological evaluation mostly differentiates tumors from hemorrhage because the margins between the tumors and hemorrhage is frequently well-defined.\textsuperscript{11,23} Intratumoral hemorrhage and tumor may be very difficult to differentiate during spontaneous Intracranial hemorrhage (ICH), even with contrast when the tumor is compressed by a large hemorrhage.\textsuperscript{11,24} While MRI offers enhanced tissue description and more accurately the location of lesions than CT-scan, both imaging procedures offer harmonizing evidence about intratumoral hemorrhage in GBM.\textsuperscript{25,26} CT-scan is a faster and more readily available instrument that offers images that are simpler to interpret than MRI in patients with GBM presenting with acute intratumoral hemorrhage. Furthermore, CT-scan is extremely helpful in evaluating the distinctive sources of T1-weighted hyperintensity on MRI images such as hemorrhage and calcification.\textsuperscript{25}

Serial MRI studies revealed delayed evolution of hemorrhage into ring enhancing lesion within a hemorrhagic GBM.\textsuperscript{6,27} This delay is probable due to the shortening of the T1 relaxation time by methemoglobin, and a magnetic predisposition consequence of intracellular hemoglobin disintegration components on the T2 relaxation time. Atlas et al observed a delay of 16 days from the hypointensity correlated to acute bleeding to the hyperintensity linked to the synthesis of uninhibited methemoglobin,\textsuperscript{27} while Destian et al. observed a lengthier delay of up to numerous months.\textsuperscript{9} We also observed a delay window of up to 14 days. The pathogenesis of this delay is uncertain so further observational studies are required to finally conclude on the delay window period. Studies have shown that methemoglobin synthesis is highest at moderately

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**Figure 3. CT-angiography showing no vascular abnormalities.**

**Figure 4. Postoperative CT-Scan images showing to total evacuation of the hematoma.**
Low oxygen tensions of approximately 20-25 mm. Subsequently, oxygen is essential for the synthesis of the mediators conscientious for the oxidation of deoxyhemoglobin to methemoglobin, besides oxygen restricting the concentration of deoxyhemoglobin availability for oxidation to methemoglobin hence the synthesis of methemoglobin is infinitesimal at zero as well as extreme oxygen tensions.

On the other hand, studies have shown that at 20-25 mm, there is sufficient oxygen to permit the synthesis of the reducing mediators, nevertheless not sufficient to diminish the concentration of deoxyhemoglobin. Therefore, analogous hypoxia inside gliomas ought to argument instead of delay the synthesis of methemoglobin. Furthermore, the delay in evolution could be due to sustained bleeding from the tumor and/or the quick absorption of hemoglobin disintegration components by an extremely vascular glioma. It’s well known that the synthesis of methemoglobin is highest at moderately low oxygen tensions, and this be the explanation for central hyper-intensity (ring enhancing lesion) normally seen on stage-2 hemorrhagic gliomas. Therefore, in a pure hematoma with very low oxygen tension at the center there is inhibition synthesis of reducing mediators obligatory for the oxidation of deoxyhemoglobin to methemoglobin and the synthesis of methemoglobin primarily arises from the periphery.

The management of newly established GBM demands a multidisciplinary method and existing standard therapy comprises of near total secure surgical resection, preceded by simultaneous radiation with temozolomide (TMZ), an oral alkylating chemotherapy drug, and then adjuvant chemotherapy with TMZ. GBMs are often invasive and affects eloquent areas of the brain. Most of these areas control speech, motor function, and the senses and so making far-reaching and total surgical resection difficult if not impossible. Furthermore, the disease succession or relapse is very common due to the extraordinary extent of invasiveness and radical surgical removal of the principal lesion is usually not beneficial since the infiltrating gliomas cells always continue to linger inside the neighboring brain. Also, modern innovations like multimodal neuronavigation and incorporation of patient-specific anatomic and functional data as well as the use of functional MRI, diffusion tensor imaging (DTI), CT-scans and MRI in preoperative planning has made surgical resection less complicated.

Notwithstanding these technical innovations above, distinguishing normal brain and residual tumor still problematic so Zhao et al. advocates the use of 5-aminolevulinic acid (3-ALA) dye as a fluorescence guide surgery. Radiotherapy and chemotherapy are very crucial components of the management of GBM patients so after surgical resection, the patient usually waits for one month for the craniotomy wound to heal before starting these other therapeutic options. Radiation therapy (RT) alone after operation was the standard management regimen until 2005, when the outcome of a pivotal phase III trial transformed this peculiar management regimen for GBM. This trial also known as the Stupp regimen demonstrated adequately that external beam of RT with simultaneous TMZ chemotherapy is more efficacious compared to RT alone.

It is worth emphasizing that multiple potential long-term hitches of whole-brain RT (WBRT) such as endocrinopathy, neurocognitive toxicity, and RT-induced leukoencephalopathy has limited the use of WBRT in the management of newly detected GBM. These grave hitches have led to further investigate into the use of involved field RT (IFRT) to minimize toxicity. Narayana indicated that IFRT dispenses external beam of RT to the tumor with a 2-3 cm perimeter with assumption that that after RT, GBM relapses within 2 cm of the primary tumor location in 80-90% of patients. Further studies have proven that the use of three-dimensional conformal beam of RT or intensity-modulated RT is the gold standard treatment and mainstream total dose of 60 Gy in 1.8-2 Gy fractions must be delivered five days per week for six weeks. Barani & Larson indicated that with RT doses to 60 Gy, patient have a more survival rate but with increased doses above this may lead to augmented toxicity with no added survival advantages.

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**Case Report**

**Figure 5. Pathological images.** A) Is the evacuated hematoma, while B-H) are images of varies immunohistochemical staining. B) HE 400×, C) GFAP 400×, D) Oligo2 400×, E) IDH-1 400×, F) ATRX 400×, G) P53 400×, H) MIB-1 400×.
The morphologically of GBM comprises of small spindle-shaped cells, characterized by polymorphism, anaplasia and distinct anisokaryosis with acidophilic cytoplasm and unclear cellular margins. They have oval or elongated nuclei with coarsely clumped hyperchromatic chromatin and multiply distinct nucleoli situated centrally or peri-centrally with a bigger nuclear to cytoplasmic ratio, nuclear pleomorphism as well as intranuclear inclusions. Furthermore, lymphocytes, neutrophils, macrophages as well as necrotic cells are also seen with very extreme vascularization. Some cells look like adipocytes because of existence of enormous lipomatous vacuoles and may make up 5-10% of all the tumor cells and as high as 80% in some individual cases. Pseudopalisades are also seen and arranged from 30-1500 μm in the greatest internal width and from 50-3500 μm in the greatest internal length with some areas showing the presence of multiple apoptotic cells. The malignancy risk of the lesion is associated with high degree of atypia, nuclear hyperchromatosis, amplified mitotic index, existence of areas of necrosis and atypical blood vessels.

OLIG2 is a transcription factor associated with synthesis of oligodendrocytes and is well noted to partake in the initiation of malignant gliomas from neural stem cells. Recently, mutations of isocitrate dehydrogenase (IDH) genes have been linked to the pathogenesis glioma. IDH-1 and IDH-2 are NADP-dependent enzymes that catalyze the synthesis of α-ketoglutarate from isocitrate during cellular metabolism. IDH-1 mutations are mostly seen concurrently with either TP53 mutations or co-deletion of 1p/19q chromosomes which means that IDH-1 mutation partake actively in the pathogenesis of infiltrating gliomas. Promoter methylation of methylguanine-DNA methyltransferase (MGMT), co-deletion of 1p/19q chromosomes, EGFR-phosphoinositide 3-kinase pathways and IDH-1 mutations have proven to be very crucial in the pathogenesis of infiltrating gliomas. Ki-67 antigen appearance indicates the percentage of cellular and biological aggressiveness in malignancy. On the other hand, p53 protein is associated with the regulation of cell cycle and has been anomalous quantification in gliomas. It’s not every intracranial glioma that is of pure vascular origin. The etiology of hemorrhage in our case although not well known is believed to be multifactorial. The management of our case was multidisciplinary and incorporated surgery, radiotherapy as well as chemotherapy.

References

Conclusions
The diagnosis of intratumoral hemorrhage in our case was very challenging at the initial stages but with time the hematoma evolved into ring enhancing images typical of GBM. The etiology of hemorrhage in our case although not well known is believed to be multifactorial. The management of our case was multidisciplinary and incorporates surgery, radiotherapy as well as chemotherapy.


