Cavernoma: New insights from an unusual case

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PII: S1878-8750(17)30436-9
DOI: 10.1016/j.wneu.2017.03.113
Reference: WNEU 5482

To appear in: World Neurosurgery

Received Date: 21 November 2016
Revised Date: 21 March 2017
Accepted Date: 23 March 2017


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Cavernoma: New insights from an unusual case

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Abstract

Rapid growth in cerebral cavernous malformation is rare. We present the case of a 71 year old patient with known multiple cavernomas over many years in whom one lesion showed rapid expansion in size. Histological examination revealed coexistence of glioblastoma within the cavernoma. We review the literature for similar cases and discuss the potential mechanisms underlying this phenomenon. Review of the literature revealed four cases with known cerebral cavernous malformations that have later developed, at the same site, a high grade glioma. All reported cases involve female patients with age ranging from 25 to 71, with imaging confirming rapid growth in the lesion. We conclude that although rare, rapid expansion of existing cavernoma should be treated as suspicious for development of other malignant tumors, and propose adding the possibility of chronic inflammation in the surrounding brain caused by micro-bleeds and hemosiderin deposition from the cavernoma to the list of possible causes.

Key Words
cavernoma, high grade glioma, glioblastoma, transformation
Introduction

Cerebral cavernous malformations (CCMs) are vascular malformations characterized by enlarged capillary cavities without intervening brain parenchyma. They are rare, with a prevalence of around 0.5% in the general population, accounting for 5-15% of all vascular malformations of the central nervous system and can be either sporadic or familial(2, 20). Familial cavernomatosis represents 10-15% of the total number of cavernoma cases and it is inherited in an autosomal dominant fashion with various degrees of penetration(8). Familial cavernomas are multiple with their risk of hemorrhage being twice as high compared to sporadic ones(17). Cavernomas are usually occult and present with an episode of symptomatic hemorrhage between the second and fifth decades of life. Other common presenting symptoms include headaches, seizures and focal neurological deficits(20).

The co-existence of cerebral cavernous malformations and primary brain gliomas is rare. It is still unknown why this phenomenon occurs with various unverified theories proposed(3, 11, 13, 18, 27). Here we present the case of a 71 year old lady who developed glioblastoma multiforme at the site of a pre-existing cavernoma. To our knowledge, there have only been five reported cases of malignant gliomas arising at the site of pre-existing cavernoma(1, 18, 21, 22, 28). Here, we present our case and discuss potential biological pathways for tumor development in patients with CCMs.
Case Report

A 71 year-old woman with a history of multiple cavernomata presented to the emergency department with a 3 days history of left sided hemiparesis, facial weakness and confusion.

In the year preceding this admission she sought the advice of her neurologist because of increased seizure frequency and intermittent, profound emotional lability. She had been diagnosed with multiple cavernomata in December 2012, after developing seizures, which were controlled with levetiracetam and lamotrigine.

Patient did not receive radiotherapy or radiosurgery for her cavernomata which were managed conservatively with imaging surveillance under the care of the neurovascular MDT (figures 1,2). She underwent formal genetic testing which identified a heterozygous single nucleotide substitution in the CCM2 gene.

On this admission, patient had left sided hemiparesis (MRC grade 4/5), facial weakness, difficulty following commands, as well as difficulty with number comprehension, calculation, and reading. MRI revealed a heterogeneously enhancing lesion with T1 shortening and peripheral contrast enhancement consistent with a high grade tumor (figure 3).

The patient underwent craniotomy and resection with histopathological examination revealing glioblastoma multiforme, WHO grade IV, IDH1 negative and cavernoma (figure 4). In the tumor sample the MGMT promoter gene was unmethylated. She was subsequently treated with radiation and chemotherapy as per Stupp protocol for the treatment of glioblastoma.
Discussion

Though rare (Table 1), the co-existence of high grade gliomas and cavernomas is recognized, but their presence in combination might be more frequent than reported. Various theories have been proposed attempting to describe the etiological relationship between the two, including (a) fortuitous association (b) sequential – secondary due to reactive or neoplastic glial change to a previously existing vascular malformation or vice versa under the influence of tumor angiogenic factor (c) genetic predisposition (d) common viral origin (e) radiation exposure (3, 11, 13, 18, 27).

The common viral origin theory has been proposed more than three decades ago but the supporting evidence is insufficient. Polyoma virus, which belongs to papovavirus family, is a pathogen only for mice and is known to have dose dependent tumorigenic effect. High doses can cause multiple capillary, cystic, or cavernous hemangiomas in brain and spinal cord with the formation of the latter greatly enhanced by suppression of the immune system (11, 25). Polyoma virus is not known to induce glial tumors but it has been shown that another papovavirus, SV 40 virus, causes ependymomas in new-born hamsters (9, 16). Although of interest, applicability of this knowledge to humans remain questionable.

We have recently gained significant knowledge surrounding the molecular genetics of cavernomas through the study of hereditary cavernoma syndromes that confirm link of the latter to three genes: CCM1, CCM2, CCM3 (6). A loss-of-function mutation to any of these genes produces an autosomal dominant syndrome that is characterized by one or more cavernomas. Cavernomas can also develop following somatic mutations of these genes (26). Interestingly, monosomy of chromosome 7 that hosts both CCM1 and CCM2, is common following hyper-activation of Ras
oncogenes or mutation of the neurofibromatosis-1 protein indicating that genetic factors within glial and Schwann cell tumors may lead to subsequent development of cavernomas (18).

Additionally, growth factors and cytokines expressed by vascular anomalies in the brain have been shown to promote tumorigenesis. Dai et al, investigated the role of platelet-derived growth factor and platelet-derived platelet factor receptor (PDGF and PDGFR) in cell cultures and in vivo in mice. They presented evidence that some low-grade gliomas may be comprised of proliferating glial progenitor cells with blocked ability to differentiate whereas malignant gliomas have acquired mutations that disrupt the cell cycle arrest pathways such as by loss of Ink4a-Arf. Co-expression of PDGF and PDGFR has been demonstrated in human gliomas indicating that autocrine stimulation may be involved in the formation of malignant gliomas. By introducing PDGF to neural progenitors and astrocytes they showed gliomagenesis with Ink4a-Arf loss promoting malignant phenotype (7). Various other factors that are secreted by gliomas or the cells surrounding them as well as vascular malformations, including vascular endothelial growth factor (VEGF) and angioprotein-1 have been shown to promote the development of cavernous malformations but also the progression of various intracranial tumors (10, 14, 23, 24).

Another possible explanation for malignant changes in glial cells adjacent to cavernomas may be chronic inflammation caused by micro-bleeds and hemosiderin deposition from the cavernoma. Chronic inflammation is known to promote malignancy in a range of tissues such as colorectal carcinoma in Chron’s disease, bladder cancer with the use of indwelling catheters and gastric adenocarcinoma with H. Pylori infection (5). It is therefore possible that similar processes over time may also lead to malignant transformation of glial cells in proximity of the cavernomas.
The fact that all cases described had a long history of prior cavernoma is consistent with such a hypothesis of chronic irritation deriving the process of tumor formation. Additionally, in three out of these five cases evidence of previous hemorrhages is reported in the form of hemosiderin deposits that were identified either intraoperatively(28) or following histopathological examination of the specimens(1, 22).

Not surprisingly, in all cases there was significant change in the size of the malformation at the time of the diagnosis of high grade gliomas, as well as new or worsening of pre-existing seizure activity with the exception of the posterior fossa case, but little is still known about the change in size of cavernomas over time. First to describe dynamic changes of these malformations was Pozzati in 1989, presenting three cases in which the cavernomas grew substantially over time(19). Similarly, rapid growth with aggressive behavior despite multiple surgical excisions in a tumor with histologic confirmation of cavernoma has been described by Stacey in 2000(12). In contradiction, Kim in 1997 showed decrease in the size over time and more specifically from an average diameter of 14.2mm to 9.1mm with repeat imaging(15). The biggest study demonstrating dynamic changes in the size of cavernomas by Clatterbuck published in 2000, did not favor either growth or shrinking over time. In this study, 68 patients with 114 cavernomas underwent serial imaging over a period of 3.7 years. 22% of the lesions were stable in size over time, 43% increased in size and 35% decreased. Interestingly, many of the above reported cavernomas had both periods of increase and decrease in size(4).
CONCLUSION

It is unusual for cavernomas to grow rapidly. We report a case with rapid expansion of a known cavernoma with histology confirming glioblastoma embedded within the malformation, radically altering the management strategy. The biological mechanism is still speculative but may be related to common genetic pathways for both lesions or tumorigenic cytokines and growth factors released by vascular malformations. We propose adding the possibility of chronic inflammation in the surrounding brain caused by micro-bleeds and hemosiderin deposition from the cavernoma to this list. From a practical point, in rapidly growing cavernomas, a low threshold to obtain histological confirmation, either in the form of biopsy or resection, should be employed to rule out malignancy.

Disclosure
There is no financial or other conflict of interests to be declared by the authors.

Patient Consent
Patient has consented to the submission of this case report to the journal.
References


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Table 1 Literature review identified five cases with malignant gliomas developing at the same site with known cavernoma. GBM: glioblastoma, CCM: cerebral cavernous malformation. RF: right frontal, LT: left temporal, LP: left parietal, V: vermis
Figure 1. Initial MRI in January 2013 showing multiple intracranial lesions with T1 shortening consistent with multiple cavernomas with evidence of previous hemorrhage
Figure 2. Interval MRI in June 2013 showing no significant changes in appearances compared to the initial MRI.
Figure 3. Imaging in April 2015 when patient presented with increasing seizure activity and hemiparesis showing significant increase in the size of the lesion with heterogeneous enhancement.
Figure 4 showing a largely necrotic pleomorphic tumor diffusely infiltrating the grey and white matter, small pieces of hippocampus and many abnormally formed blood vessels. The tumor is composed of large cells with pleomorphic oval-shaped nuclei and prominent nucleoli arranged in solid bundles within a fibrillary background. Large areas of pseudopalisading necrosis, endothelial cell proliferation with glomeruloid structure formation and brisk mitotic activity is noted. There are areas with collection of hyalinised and dilated blood vessels with different caliber and wall thickness; some with thrombosis and calcification. EVG confirms the vascular malformation consistent with cavernoma.
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Highlights

- Five reported cases with co-existence / development of high grade gliomas in patients with long history of cavernomas
- Rapid growth in a cavernoma should raise suspicion of malignancy
- Theories proposed cannot explain co-existence of two lesions
- We propose adding chronic inflammation due to micro-bleeds as causative mechanism
Cavernoma as a benign vascular malformation: is it time to rethink?

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**Abbreviations**

CCM: cerebral cavernous malformation
MDT: multi-disciplinary team
WHO: World Health Organisation
MRC: Medical research council
MRI: magnetic resonance imaging
IDH: isocitrate dehydrogenase
MGMT: O⁶-methylguanine-DNA-methyltransferase
PDGF: platelet-derived growth factor
PDGFR: platelet-derived platelet factor receptor