Supratentorial cerebral cavernous malformations: clinical, surgical, and genetic involvement

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Object. Although there is general agreement on the methods of treatment for symptomatic supratentorial cerebral cavernous malformations (CMs) located in noneloquent areas, some controversy exists regarding the management of cerebral CMs that are asymptomatic and located in eloquent or deep areas. Moreover, recent advances in genetic findings could influence both standard clinical management and the follow-up strategy in affected individuals. Thus, the objective of this study was to develop, based on the authors’ experience and a literature review, a management algorithm to deal with supratentorial cerebral CMs.

Methods. The authors retrospectively reviewed the clinical data related to 118 patients who underwent surgery for symptomatic supratentorial cerebral CMs at their institution. Twenty-eight of 118 patients harbored multiple lesions, and nine of these 28 patients had a clinically positive familial history. Genetic investigations were performed in 89 patients (75%).

Conclusions. Surgery for supratentorial cerebral CMs in noneloquent locations is safe and curative. In cerebral CMs located in deep and eloquent areas and with symptoms including progressive neurological deficits, evidence of hemorrhage, and uncontrolled seizures, surgical treatment according to an integrated plan based on frameless stereotactic guidance and functional magnetic resonance imaging is recommended and results in acceptably low morbidity. The data support the need for long-term imaging follow up in all patients, careful preoperative vascular studies to detect associated venous anomalies, and the importance of genetic mutational analysis. The DNA screening protocol will change the care of family members of patients with familial forms of cerebral CMs, because affected asymptomatic family members may benefit by early detection of lesions. At the same time, the exclusion of family members who are not carriers of the mutation as members of the population at risk reduces the economic and psychological burden of clinical and instrumental monitoring.

Key Words • cerebral cavernoma • cavernous angioma • management algorithm • hemorrhage • vascular malformation

Cerebral CMs are angiographically cryptic lesions that are classically defined by enlarged and thinned walled vascular structures in the central nervous system without intervening brain parenchyma. These malformations are lined by endothelial cells and lack supporting vascular smooth-muscle cells, and they are surrounded by hemosiderin deposits and gliosis, which may or may not be thrombosed. On gross examination, they appear as berry-like, reddish-purple lesions that are variable in size (ranging from 1 mm to several centimeters), multiple or single, often encapsulated and multilobar, containing hemorrhage in various stages of organization, and occasionally calcified.

The true prevalence of cerebral CMs in the general population is unknown. These lesions have been found in up to 0.5% of the population in large autopsy and MR imaging series. The clinical presentation of these lesions is highly variable, ranging from an incidental finding at neuroimaging to discovery during autopsy after a fatal hemorrhage. The most common symptoms of CMs are seizure disorders (possibly caused by the toxic effects of iron deposition), followed by focal neurological deficits and headache.

In virtually all CMs, there are signs of repeated microhemorrhage on neuroimaging, even if clinically significant hemorrhage is a less common phenomenon (with an annual risk at 0.25 to 6%), and the incidence of microhemorrhages increases after the first episode. Nevertheless, this level of risk, if we consider it over a lifetime in young patients and in the few cases in which massive fatal hemorrhage has occurred, is not insignificant. The MR imaging modality is the diagnostic tool of choice because of its high sensitivity and specificity in detecting the hemoglobin degradation products present inside and around the lesion.

Cerebral CMs can be found in every region of the cen-
tral nervous system, and the supratentorial location is represented in almost 70 to 80% of intracranial cases, with multiple lesions in 10 to 20% of patients. Although cerebral CMs were once considered to be a developmental disorder, several authors have reported their de novo appearance, mostly after radiation therapy.4,10,15,27,34,56,62 Cerebral cavernous angiomas can occur in a sporadic or autosomal-dominant inherited form, with familial cases often characterized by the presence of multiple lesions, whereas patients in nonfamilial cases usually harbor only a single malformation.25,48

Although there is general agreement on the method of treatment for symptomatic supratentorial cerebral CMs located in noneloquent areas, some controversy exists about the management of asymptomatic lesions and malformations located in eloquent or deep areas. Moreover, recent advances in genetic findings could influence both standard clinical management and the follow-up strategy in affected individuals.25,48 Thus, the goal of this study was to develop, based on our experience and a literature review, a management algorithm to deal with supratentorial cerebral CMs.

Clinical Material and Methods

Patient Population

We retrospectively reviewed the clinical data related to 118 patients who underwent surgery for symptomatic supratentorial cerebral CMs at our institution between January 1992 and August 2005. The patients’ ages ranged from 19 months to 90 years (mean 39 years), and the male/female ratio was 1.8:1. The mean follow-up duration was 55 months (range 8–164 months).

Number and Characteristics of Lesions

We removed 125 cerebral CMs: 90 in patients with single lesions and 35 in a group of 28 patients with multiple cavernomas. In this latter group, nine patients had a positive familial history for cerebral CMs.

The size of the cavernomas ranged from 4 to 60 mm (mean 26 mm). One hundred ten of 125 cerebral CMs were deep seated (lateral ventricles, corpus callosum, thalamus, basal ganglia).

Clinical Presentation

The main clinical presentation included epilepsy (69 patients), focal neurological deficits (37), raised intracranial pressure (six), and chronic headaches (six). Clinical symptoms related to hemorrhage were acute or subacute in 43 cases and chronic in six. Hemorrhage, which was present in 38% of patients, was intraleisional in six cases, extralesional in 35, and intraextralesional in two. In two other patients, subarachnoid hemorrhage was observed in association with an extralesional hemorrhage. The size of the hemorrhage was less than 2 cm in 25 lesions and more than 2 cm in 20. Six cavernomas contained a capsulated hemorrhage.

Seizures, which were present in 69 (58%) of the patients, were of the following types: generalized tonic clonic in 10 patients (14%), generalized tonic clonic with focal onset in 11 (16%), simple partial type in 28 (41%), and complex partial in 20 (29%). The duration of epilepsy history was less than 6 months in 45 patients (65%) and more than 6 months in 24 (35%). The Engel classification17 (Table 1) was used to evaluate postoperative seizure outcome.

Neuroimaging Studies

A 3-tesla fMR imaging unit was used in 10 patients for motor strip identification, whereas cerebral digital subtraction angiography was performed for identification of associated vascular anomalies in 56 patients with hemorrhage or lesions located in deep or eloquent areas.

Surgical Procedures

Surgery was performed within 10 days in patients presenting with acute intraextrallesional hemorrhage. The surgical approach was transcortical in 102 cavernomas, transsulcal in 17, and interhemispheric–parasagittal in six. In 39 patients with subcortical, central region, or deep-seated cavernomas, intraoperative anatomical guidance was provided with the aid of a neuronavigational system (VectorVision; BrainLAB USA, Inc., Moorestone, NJ). Intraoperative neurophysiological mapping was performed in 10 patients and integrated fMR imaging with neuronavigation in five, all with lesions located in eloquent areas.

Genetic Studies

Genetic studies were performed in 89 patients (75%), as follows: the group of nine patients with a family history of cerebral CMs, all of whom were harboring multiple lesions; a group of 18 patients harboring multiple lesions in whom there was no family history of cerebral CMs; and a group of 62 patients with single, apparently sporadic lesions, but in whom the family history either was not complete or was not suggestive of possible cerebral CMs (that is, seizures, cerebral hemorrhage of unknown cause, and so on).

Statistical Analysis

The data were statistically analyzed using the chi-square test. The Fisher exact probability test was used when there were fewer than four variables. The level of statistical significance was set below 0.05.

Results

No significant difference in the incidence of CMs was found among the various age groups or according to sex. Developmental venous anomalies were identified on angiographic studies in 41 (73%) of 56 patients. The anomalies designated as venous angioma and distinguished from angiomas that were “venous-like” were present in 19 patients (34%), with 12 anomalies adjacent (Fig. 1) and seven distant to the cavernoma (Fig. 2). Other venous anomalies included hypertrophic veins, thrombosed veins, and hypoplastic or thrombosed sinus (present in 22 cases [39%], with 11 anomalies adjacent and 11 distant to the cavernoma); capillary blush in three cases (5%), arterial afferent in two cases (3%), other vascular lesions in two cases (3%).
cases (3%), one arteriovenous malformation, and one aneurysm.

Radical lesionectomy was performed in 124 lesions (99%), with a one-stage double lesionectomy in seven patients. Partial resection was performed in one cavernoma located in the thalamus. A gliotic hemosiderin ring was totally resected in 61 cerebral CMs, all of them located in cortical–subcortical noneloquent areas.

There were no surgery-related deaths in this series. In 78 patients (66%) the neurological condition remained unchanged after surgery, whereas 30 patients (25%) presented with clinical improvement. The neurological condition deteriorated in three patients (2.5%) after surgery, all with postoperative complications: one patient had an intraparenchymal hematoma secondary to removal of an associated venous angioma, which required no treatment; one patient who underwent evacuation of hematoma from the residual cavernoma; and one who underwent removal of a basal ganglia cerebral CM and who suffered from postoperative ischemia at the site of surgery. Other postoperative complications were as follows: one extradural hematoma, which required no treatment; one case of pulmonary embolism; and two cases of deep venous thrombosis. The patients remained in the intensive care unit for 1 to 6 days after surgery, and the mean hospital stay was 7 days.

No neurological deficit was found at follow-up evaluation in 80% of patients, a moderate neurological deficit was found in 13%, and severe deficit in 7%. Patients with moderate and severe disability had lesions located in eloquent or deep cerebral areas.

At follow-up examination in the 69 patients suffering from preoperative seizure disorders, 57 (82%) were in Class I, six (9%) were in Class II, four (6%) were in Class III, and two (3%) were in Class IV, according to the Engel classification (Table 1), with a significant correlation between a seizure history shorter than 6 months and better outcome after epilepsy surgery (Table 2). The correlation between simple lesionectomy with or without removal of the hemosiderin gliotic ring and postoperative outcome in patients with epilepsy was not significant. A second-stage additional amygdalohippocampectomy was performed in one patient for uncontrolled seizures, with subsequent control. On the last follow-up imaging session, two recurrences were revealed adjacent to the resected cavernoma.

In 13 patients, a de novo development of cerebral CMs distant from the initial lesion was discovered: in 12 of 13 patients from the group with clinically and/or genetically confirmed familial cerebral CMs, and in one of 10 patients from the group with sporadic lesions. Genetic studies resulted in the following findings: seven mutations (78%) in the group of nine patients with a positive family history and multiple cerebral CMs; 12 mutations (66%) in a group of 18 patients harboring multiple lesions with no family history of cerebral CMs; and two mutations (3%) in the group of 62 patients with a single, apparently sporadic lesion. To date, DNA analysis in family members...

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<tr>
<td>I</td>
<td>free of disabling seizures</td>
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<tr>
<td>II</td>
<td>rare disabling seizures, “almost seizure free”</td>
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<td>IV</td>
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* According to the scale published by Engel, et al., in 1993.

**TABLE 1**

Engel classification

**ENGEL CLASSIFICATION**

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**Table 1**

**Figure 1.** Left: Axial MR image showing a right temporal cavernous angioma. Right: Angiographic study demonstrating an associated adjacent venous angioma-like lesion (arrows).
has shown 40 carriers of the mutation, 19 of them harboring asymptomatic cerebral CMs at the time of MR imaging examination.

Discussion

There is general agreement in recommending surgery for symptomatic supratentorial cerebral CMs that are not located in deep or eloquent areas. In these cases, surgery is generally safe and its efficacy is established both for control of epilepsy and prevention of rebleeding, whereas controversy exists about resection of cerebral CMs located in critical areas in patients with few or no symptoms. In agreement with other authors, we found that supratentorial cerebral CMs, including those in critical areas, can be microsurgically treated with a favorable risk/benefit ratio by using, on a case-by-case basis, frameless stereotactic guidance, electrophysiological monitoring, and/or integrated neuronavigation with fMR imaging.

The epileptogenicity of cavernomas has been ascribed to the ongoing deposition of iron and blood breakdown products. We were unable to detect a significant relationship between a postoperative hemosiderin ring and seizure outcome. In our series, lesionectomy alone with or without resection of the gliotic hemosiderin ring accomplished good seizure control in the majority of patients with epilepsy, and there was a significant correlation of good outcome with a shorter seizure history (Table 2), in agreement with other authors. In some cases of mesial temporal cavernomas with chronic or partially controlled seizures, the use of intraoperative electrocorticography may help us decide whether to perform further resection—that is, amygdalohippocampectomy or standard anterior temporal lobectomy. We performed additional amygdalohippocampectomy in a second stage in a patient with poor seizure control after a complete pure cavernoma resection. As reported and in agreement with our experience, resection of associated venous anomalies contiguous to the cavernoma could be dangerous and should be avoided.

In our series, routine MR imaging follow-up studies revealed two recurrences of the resected cerebral CMs in adjacent locations and 13 distant de novo cerebral CMs, mostly in familial forms. None of these patients had undergone radiation therapy, nor had they suffered viral infections during the follow-up period. In agreement with other authors, we believe that this discovery may lead to an upward revision in risk estimates, because hemorrhage risk per annum has traditionally been calculated since birth. Therefore, we emphasize the need for long-term imaging follow up, mostly in familial cases.

Genetic factors have been identified in 20 to 30% of patients with CMs. Familial cases are often characterized by the presence of multiple lesions, whereas sporadic cases usually entail only a single malformation. In familial cases, the clinical symptoms denoting penetrance are not always present, and it has been suggested that this might depend on the gene causing the cerebral CMs and on the patient’s age at onset. However, in other reports, it has been suggested that up to 75% of the so-called sporadic cases with multiple lesions are, in reality, familial cases, with asymptomatic vascular lesions in relatives masking the autosomal-dominant segregation pattern. Based on MR screening studies, in familial cases, penetrance is age-dependent and approaches 100% in affected adults, even if only approximately 60% of those affected

<table>
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<tr>
<th>Postop Engel Class</th>
<th>Preop Seizure History*</th>
<th>No. (%) w/ ≤ 6 Mos</th>
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<tr>
<td>I</td>
<td>41 (91)</td>
<td>16 (66)</td>
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<tr>
<td>II</td>
<td>3 (7)</td>
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<td>IV</td>
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* In 45 patients the preoperative duration of seizure history was 6 months or less, and in 24 patients it was more than 6 months.

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Fig. 2. Left: Axial MR image showing a left rolandic cavernous angioma. Right: Angiographic study demonstrating an associated distant venous angioma-like lesion (arrow).
Supratentorial cerebral cavernous malformations

have clinically recognizable symptoms.25,32,49,58 This sup-
ports the suggestion that the population incidence of fa-
miliar cerebral CMs may be significantly underestimated.

In our experience, genetic studies were helpful in iden-
tifying mutations in seven (78%) of nine patients with a
family history of cerebral CMs; in 12 (66%) of 18 patients
harboring multiple cerebral CMs but with no known clini-
cally affected relative; and, interestingly, in two (3%) of
62 patients harboring a single, apparently sporadic lesion.
Performing mutational analysis in family members, we
found 40 mutation carriers and, among them, 19 with
asymptomatic cerebral CMs. Our data show that mutation
analysis is of striking importance for prognosis and clin-
cal management of both apparently sporadic and familial
cerebral CMs. The DNA screening protocol will change
the care of familial forms of cerebral CMs, because affect-
ed asymptomatic family members may benefit by early
detection of lesions, whereas the exclusion of nonmuta-
tion-carrying family members as at-risk individuals
reduces the economic and psychological burden of clinici-
and instrumental monitoring. In relatives of patients
with known familial forms of cerebral CM in whom the
mutations have not been found, all family members at risk
have to be monitored.

After analysis of this series of patients and a literature
review,21,29,35,38,39,54,57,60,61 we developed a management algo-

tithm to deal with supratentorial cavernomas, taking into
account clinical presentation, lesion location, and genetic
findings. The algorithm is as follows.

1. Asymptomatic cerebral CMs are generally observed
carefully, with follow-up MR imaging performed at year-
ly or 2-year intervals.

2. Symptomatic (severe headache, seizures, progressive
neurological deficits) cerebral CMs in noneloquent areas
should be resected with the aid of frameless stereotactic
guidance. In eloquent and/or deep locations, symptomatic
(intractable seizures, severe or repeated hemorrhage, pro-
gressive neurological deficits) cerebral CMs should be
resected with the aid of frameless stereotaxy and integrat-
ed fMR imaging studies.

3. In cases of cerebral CMs presenting with seizure, the
threshold for intervention depends on lesion accessibility,
eloquent location, and severity of the seizure disorder as
well as drug resistance. Patients with multiple CMs should
be studied extensively to decide if one or more lesions are
responsible for the symptoms. In noneloquent and acces-
sible areas, surgery should be performed to remove the
lesion as soon as possible after the seizure disorder begins.
In cases of temporal lobe seizures, if simple lesionectomy
tails to correct the disorder, detailed cortical and electrode
electroencephalographic mapping should be performed,
possibly followed by epilepsy surgery (such as amygdalo-
hippocampectomy).

4. Associated venous anomalies should be spared dur-
ing surgery for cerebral CMs.

5. In suspected familial cases identified based on clini-
cal and/or neuroimaging results, genetic analysis must be
performed. In the event of positive findings on mutation-
al analysis, the genetic study should be extended to family
members, who could benefit in two ways: 1) in the event that a
mutation carrier is found, by early detection of
cerebral CMs; and 2) in the event that no mutation is
found, by exclusion of the family member from the at-risk
group. Clinical and neuroimaging monitoring should be
recommended also in cases of negative findings on mu-
tational analysis but suspicious clinical symptoms or a
known family history of the disorder. When the family
history is not available or is unreliable, genetic analysis
should also be considered in patients harboring an appar-
ently single, sporadic lesion to detect cryptic or de novo
genetic mutation.

6. In cases of incomplete resection, repeated surgery is
advocated for cerebral CMs located in noneloquent areas,
whereas in a deep and/or eloquent location, further sur-
gery should be performed on a case-by-case basis.

7. We support the need for long-term follow-up imag-
ing in all patients to detect recurrences or de novo lesions.

Conclusions

Surgery for supratentorial cerebral CMs in locations
that are not eloquent or deep is generally safe and curative.
In cerebral CMs located in deep and/or eloquent areas and
accompanied by progressive neurological deficits, evi-
dence of hemorrhage, and uncontrolled seizures, surgery
performed with the aid of frameless stereotactic guidance
and fMR imaging according to an integrated plan is rec-
ommended and is followed by acceptably low morbidity.
Our data support the need for long-term imaging follow
up in all patients, careful preoperative vascular study to
detect associated venous anomalies, and the importance
of genetic mutational analysis.

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