New nidus formation adjacent to the target site of an arteriovenous malformation treated by Gamma Knife surgery

Report of 3 cases

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New nidi are rarely found adjacent to the resection margin following treatment for an arteriovenous malformation (AVM), especially in adults. In addition, there are no reports in adults of new nidus formation adjacent to the targeted site of an AVM that angiography has verified to be completely obliterated by radiosurgery. The authors present their experience with recurrent AVMs following AVM radiosurgery in 3 patients whose ages were 9 years, 10 years, and 33 years. None of the patients had been treated with embolization before radiosurgery. Two patients had a history of intracerebral hemorrhage before radiosurgery. New lesions developed around the obliterated nidi in all 3 cases. Angiography performed after the first radiosurgery confirmed complete removal of the nidi in all 3 patients, and new nidus formation was detected 31, 132, and 36 months after the initial GKS. The new lesions were also treated by GKS. Occasionally, in patients with recurrent AVMs, such as those described in this paper, long-term clinical and angiographic follow-up may be required, even if complete occlusion is originally shown on angiograms.

Case Reports

Case 1

The first patient was a 9-year-old boy treated by GKS for a left occipital AVM 1 month after he experienced ICH. The treated volume was 600 cm³, and 25 Gy was directed to the margin of the nidus. Follow-up angiography performed 1 year after radiosurgery showed a decreased size of the nidus at the same location. Thirty-one months after the initial GKS, follow-up angiography showed that the targeted nidus had been completely obliterated, but new nidus formation was noted inferior to the earlier lesion. We therefore performed a second GKS to treat this new nidus, and the cumulative treated volume increased to 900 cm³ (Fig. 1). On cerebral angiographic studies, the new lesion completely disappeared 2 years after the second radiosurgery. Meticulous clinical follow-up examinations continue to be performed in this patient for the detection of an asymptomatic AVM recurrence.
New nidus formation

Case 2

The second patient was a 10-year-old boy treated by GKS for a small AVM located in the cerebellar vermis 1 month after he initially presented with bleeding. The treated volume was 700 cm³ (Fig. 2A and Fig. 3A and D). Five years after radiosurgery, the patient suffered from repeated hemorrhage at the same site, and angiography revealed a partially occluded nidus with a total volume of 400 cm³. The patient underwent GKS 2 months after the repeated hemorrhage with the radiation dose directed to the same nidus (Fig. 2B and Fig. 3B and E). Six years after the second treatment, routine MR imaging revealed a new lesion located medial to the previously targeted nidus, which was finally completely occluded (Fig. 2C). Follow-up angiography showed the volume of this new nidus to be 2800 cm³ (Fig. 3C and F). A third GKS was successfully performed to treat the new nidus. Two years later, MR angiography confirmed that the new lesion had been completely obliterated (Fig. 2D). We plan to perform an additional angiographic study 5 years after the last radiosurgery to check for an asymptomatic recurrent AVM.

Case 3

The third patient was a 33-year-old man treated with GKS for an incidentally detected cerebellar AVM whose nidus volume was 1100 cm³. A radiation dose of 20 Gy was delivered to the lesion’s margin (Fig. 4A and B). Three years after radiosurgery, the patient experienced a cerebellar ICH and angiography revealed a new nidus adjacent to the previously targeted lesion, which was completely occluded (Fig. 4C and D). Radiosurgery was repeated for this new lesion whose volume was 500 cm³. Unfortunately, we do not have additional information on the patient’s response to the last GKS because he was lost to follow-up.

Discussion

Literature Review

Curative treatment of cerebral AVMs can be performed in a variety of ways: complete resection of abnormal vasculature, such as the nidus and arteriovenous shunting, by using a microsurgical procedure; total obliteration of the nidus by radiosurgery; or total embolization of the nidus using an endovascular procedure. Angiographic evidence of an absent nidus following these treatments generally indicates no further risk of hemorrhage; therefore, additional angiographic studies are not necessary. Recurrence after the disappearance of an AVM nidus has been reported occasionally, however, with 30 cases reported in the English-language literature. Nevertheless only one pediatric case of recurrent AVM after radiosurgery has been reported. In this article, we present three cases of AVM recurrence after successful GKS—one in an adult and two in children (Table 1). This is the first report of AVM recurrence in an adult after complete obliteration of the nidus following radiosurgery.

Kader et al. reported 5 pediatric cases of AVM recurrences in patients in whom postoperative angiograms yielded negative findings; in those cases, the recurrences appeared 1 to 9 years after the initial resection. Similarly, Klimo et al. described 5 pediatric cases of recurrent AVMs; in all these cases postoperative angiography had...
confirmed that the original lesion had been completely resected. The authors reported that in all cases the AVM recurred within 6 years, including 1 patient who experienced 2 recurrences, and that the diffuse type of AVM was more likely to recur.

Although AVM recurrences after successful surgical or radiosurgical treatment have been reported in children, these events remain extremely rare in the adult population. Previously, only 5 adult cases of recurrent AVM with negative results on postoperative angiograms were reported in the literature.4,7,8,13, 25 Gabriel et al.8 first reported an AVM recurrence in an adult after complete resection. The patient was a 19-year-old man with angiographic confirmation of complete AVM excision; recurrence occurred 9 years after the initial operation. The remaining four adults with AVM recurrences included 3 men and 1 woman ranging in age from 21 and 33 years. In these patients, the latency period between initial resection of the lesion and recurrence ranged from 4 to 9 years. Codd et al.2 presented the first adult case of double recurrence of an AVM in a 33-year-old woman with a Spezler-Martin Grade III AVM in the left occipital lobe. She experienced 2 recurrences, with the first occurring 7 years after initial complete resection and the second detected 2 years after the second resection. As an alternative treatment, stereotactic radiosurgery with an 18-Gy dose was performed.

We believe that we are the first to report an adult case in which the patient underwent radiosurgery, received confirmation of total obliteration of the nidus, and then experienced an AVM recurrence. We are reporting similar experiences in pediatric cases, but an AVM recurrence after radiosurgery has already been reported in a pediatric patient. Rodríguez-Arias et al.24 presented the case of a 9-year-old patient in whom an AVM recurred in a different location 2 years after GKS had eradicated the lesion with a 25-Gy dose to the 75% isodose line. The authors suggested that the redistribution of blood flow after progressive occlusion of the AVM may have acted on immature vessels surrounding the malformation, giving rise to another AVM. These authors also proposed that repeated angiography at the beginning of adult life should be considered for successful and consistent exclusion of any recurrence of AVMs treated by radiosurgery in pediatric patients.

**Pathogenesis of Recurrent AVMs After Operative Resection.** Several reports in the literature suggest possible theories for the pathogenesis of recurrent AVMs after complete resection. Gabriel et al.8 proposed that a small residual nidi may be related to an AVM recurrence. In that case, these recurrences may not be true recurrences, but could trigger events that could conceal the nidus temporarily on angiographic studies. Factors associated with residual nidi, such as vessel spasm, brain swelling with a compressive mass effect, or temporary thrombosis of residual AVM vasculature in the immediate postoperative period, could explain the absence of the residual nidi on angiograms obtained immediately postoperatively.

Also, small immature AVM vessels, which are generally detected in pediatric cases, may be a potential factor in AVM regrowth.9 Although these vessels may appear invisible on angiograms, they may retain the ability to form new AVMs in the same location. Likewise, Padget21 presented a fistula model consisting of an immature aberrant vessel connection between the primitive arteries and relatively large veins occurring during childhood. It has been further hypothesized that reduced resistance to blood flow in such arteriovenous fistulas may cause abnormal arteries to dilate adjacent to normal arteries.

Another alternative theory is related to the angiogenesis stimulated by brain injury, tumor, ischemia, and/or inflammation. Schmit et al.27 hypothesized that localized ischemia and inflammation could induce AVM recurrence by hyperstimulating angiogenic factors, as suggested in a de novo AVM model in an 11-year-old patient with moyamoya. Sonstein et al.28 proposed that VEGF expression may be associated with regrowth of AVMs in both pediatric and adult patients. Furthermore, Kader et al.16 suggested that surgery itself, performed during an active growth stage in patients in whom proangiogenic tissue remains after resection, may produce an angiogenic process, potentially increasing the risk of recurrence. They also suggested that VEGF may play a role in the recurrence of AVMs. Previous reports by Hashimoto et al.11 showed that AVM vessels in pediatric patients had a tendency to have a high Ki-67 index. These results offer evidence of increased endothelial cell turnover in AVMs, which may be indicative of an active angiogenic process.

Another possible hypothesis to explain true AVM

**Fig. 2.** Case 2. Axial T2-weighted MR images obtained 1 year (A), 5 years (B), 11 years (C), and 13 years (D) after radiosurgery. One year after radiosurgery, the nidus is still near the left side of the fourth ventricle (A). Five years after radiosurgery, the nidus remains at the same location, and a second radiosurgery is performed to treat this partially occluded nidus (B). Six years after the second radiosurgery, a new nidus is now observed at the midline vermis (C). A third GKS was successfully performed to treat the new nidus. Two years later, MRI confirmed that the new lesion had been completely obliterated (D).
recurrence is the “hidden compartment” model. This concept suggests that a change in the vascular supply, as shown by differences in the feeding artery between the initial and recurrent AVM, could indicate the presence of multiple compartments. Each compartment could have feeding arteries, draining veins, and nidi that are separate from the excised unfilled compartment. Although unseen on early postoperative angiographic studies, the partially thrombosed sector may have a vessel that connects with vessels in other compartments. Similar to this hypothesis, there is the “reserve nidus” concept suggested by Sano et al., which is defined as an abnormal vascular group adjacent to the primary nidus, which subsequently grows to become a recurrent lesion capable of rebleeding.

Possible Pathogenesis of Recurrent AVMs in Radiosurgery Cases in This Series. These aforementioned hypotheses could be applied to the recurrence of an AVM after total occlusion of the nidus following radiosurgery, such as that seen in the present cases. Although we cannot fully explain the factors related to AVM recurrence after radiosurgery, we can consider that the cause of the recurrence is the redistribution of blood flow after progressive occlusion of the AVM, as suggested by Rodriguez-Arias et al. In addition, we cannot completely exclude a new theory that suggests that radiation can induce angiogenesis of the surrounding nidus, in contrast to the previous concept that radiation inhibits angiogenesis of the AVM nidus. This theory is supported by the present study, in which all three patients had recurrence adjacent to the targeted site of the AVM. The lesion was located in the previous margin area and could have been affected by the radiation effect of the previous radiosurgery.

In our first case, the patient was treated with radiosurgery 1 month after experiencing hemorrhage. Because the new nidus was noted adjacent to the previous nidus, we could hypothesize that there was reexpansion of a hidden compartment.
compartment compressed by a hematoma. However, on the follow-up angiograms obtained 1 year after radiosurgery, the nidus volume was smaller and no new lesion was detected around the targeted nidus. Then 31 months after radiosurgery, a new nidus appeared adjacent to the targeted lesion, which had been completely occluded. In this case, the theory of reexpansion of a residual compartment, such as a nidus compressed by hematoma, can be discarded and the theory of a true AVM recurrence may apply. In the second case, a newly enlarged AVM was noted adjacent to the previous nidus after 2 rounds of radiosurgery. After the first radiosurgery, the patient experienced rebleeding at the same location due to partial occlusion of the AVM. After the second radiosurgery, the patient exhibited no symptoms, but a new nidus was noted 11 years after the first radiosurgery, with a volume larger than that of the initial nidus. The possibility of reexpansion of the nidus that had been compressed by a hematoma can be discarded because of the long-term follow-up interval. Also, the significantly larger volume of the new AVM, compared with that of the previous target volume, does not fit into the theory of residual nidus expansion. In the third case, the patient had no ICH at the initial radiosurgery, and the targeted nidus was completely occluded after radiosurgery. However, ICH inferior to the previous target developed 3 years after radiosurgery. This case also presents as the development of a new nidus, because there was no possibility of reexpansion of the compressed nidus.

On the basis of our analysis, we believe that our three cases are “true” AVM recurrences for several reasons. First, postradiosurgery angiograms demonstrated no obvious spasm and no thrombosis that potentially could have concealed a residual nidus. Second, the time interval between radiosurgery and follow-up angiography was long enough to recover from the pathological conditions mentioned earlier. Finally, we did not perform embolization for these AVMs, which could have led to recanalization of the initial nidus.

Conclusions

The present cases suggest that a new nidus can be found adjacent to the target site of an AVM after GKS. In addition, we describe the first adult case of AVM recurrence after complete occlusion was verified by post-radiosurgery angiography. Our cases document that angiographic confirmation of total absence of the nidus does not always eliminate the risk of AVM regeneration with the potential of rebleeding, even in adults. Therefore, some patients whose AVMs have been eliminated following resection or radiosurgery, such as the patients presented in this paper, may require long-term clinical and angiographic follow-up.

Disclosure

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

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