Targeted intraarterial anti-VEGF therapy for medically refractory radiation necrosis in the brain

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Radiation necrosis (RN) is a serious complication that can occur in up to 10% of brain radiotherapy cases, with the incidence dependent on both dose and brain location. Available medical treatment for RN includes steroids, vitamin E, pentoxifylline, and hyperbaric oxygen. In a significant number of patients, however, RN is medically refractory, and the patients experience progressive neurological decline, disabling headaches, and decreased quality of life.

Vascular endothelial growth factor (VEGF) is a known mediator of cerebral edema in RN. Recent reports have shown successful treatment of RN with intravenous bevacizumab, a monoclonal antibody for VEGF. Bevacizumab, however, is associated with significant systemic complications including sinus thrombosis, pulmonary embolus, gastrointestinal tract perforation, wound dehiscence, and severe hypertension. Using lower drug doses may decrease systemic exposure and reduce complication rates. By using an intraarterial route for drug administration following blood-brain barrier disruption (BBBD), the authors aim to lower the bevacizumab dose while increasing target delivery.

In the present report, the authors present the cases of 2 pediatric patients with cerebral arteriovenous malformations, who presented with medically intractable RN following stereotactic radiosurgery. They received a single intraarterial infusion of 2.5 mg/kg bevacizumab after hyperosmotic BBBD.

At mean follow-up duration of 8.5 months, the patients had significant and durable clinical and radiographic response. Both patients experienced resolution of their previously intractable headaches and reversal of cushingoid features as they were successfully weaned off steroids. One of the patients regained significant motor strength. There was an associated greater than 70% reduction in cerebral edema.

Intraarterial administration of a single low dose of bevacizumab after BBBD was safe and resulted in durable clinical and radiographic improvements at concentrations well below those required for the typical systemic intravenous route. Advantages over the intravenous route may include higher concentration of drug delivery to the affected brain, decreased systemic toxicity, and a significantly lower cost.

http://thejns.org/doi/abs/10.3171/2014.9.PEDS14198

KEY WORDS bevacizumab; radiation necrosis; radiation adverse effect; blood-brain barrier disruption; arteriovenous malformation; stereotactic radiosurgery; intraarterial chemotherapy; oncology
Providing a rationale for further study. By using an intravenous route for drug administration following blood-brain barrier (BBB) disruption (BBBD), we aimed to lower the bevacizumab dose while increasing target delivery.

**Case Reports**

This case report was reviewed by the University of Louisville institutional review board.

**Technique for Intraarterial Bevacizumab Infusion After BBB Breakdown**

The patients were premedicated with 6 mg dexamethasone and 1000 mg Keppra. The femoral artery was accessed using the Seldinger technique. A 5-Fr diagnostic catheter was used to catheterize the cervical internal carotid artery ipsilateral to the lesion. Baseline internal carotid artery angiography was performed.

Next, 5 mg Valium and 0.2 mg atropine were administered intravenously. Warm (37°C) 25% mannitol was infused into the ipsilateral cervical carotid artery at a rate of 4 ml/sec for 30 sec. This was followed by infusion of 2.5 mg/kg bevacizumab in a volume of 100 ml over 14 minutes.

The patients were then monitored in the recovery unit with frequent neurological exams and continuous heart rate and pulse-oximetry measurement for 2 hours. They were then observed overnight in a pediatric intensive care unit, with neurological status and vital signs checked every 2 hours, as well as continuous heart rate and pulse-oximetry monitoring. Both patients were discharged to home on postoperative Day 1. No adverse events were noted.

**MRI Signal Quantification**

The MRI scans were retrospectively reviewed by a board-certified neuroradiologist, and the volume of altered FLAIR signal and Gd enhancement was analyzed. The images were viewed on an Aquarius workstation (TeraRecon). Utilizing the segmentation, analysis, and tracking module, the regions of interest were outlined and volumetric data were calculated. The follow-up data were compared with the baseline data to produce a calculation of percentage change from baseline.

**Case 1**

**History and Examination**

A 12-year-old right-handed girl presented with severe headaches. Brain MRI showed a 3.2-cm left posterior frontal arteriovenous malformation (AVM; Spetzler-Martin Grade III) and mild surrounding vasogenic edema. There were no signs of hemorrhage. The patient was initially started on 24 mg per day of dexamethasone that was tapered over 3 weeks. Because of worsening headache and progressively worsening right hemiparesis, repeated attempts to reduce dexamethasone dose below 6 mg daily proved unsuccessful over the ensuing 9 months. Courses of pentoxifylline, vitamin E, and hyperbaric oxygen were all administered per study protocols, without success. Despite these interventions, her proximal right upper and lower extremities declined to Grade 4/5 strength and her right hand and foot declined to Grade 0/5 strength. She experienced a 60-pound weight gain and severe cushingoid features. She complained of constant severe headache (10/10 on the visual analog scale). She also experienced severe emotional lability, causing her to withdraw from school.

**Intervention**

Given the progressive neurological decline and the lack of alternative therapies, the option of intravenous administration of bevacizumab was initially explored. However, this proved to be not available for our pediatric patient population. Hence, we considered the option of low-dose intraarterial bevacizumab treatment. Our rationale for this approach was to use a smaller dose of bevacizumab in a more directed fashion (intraarterial administration after disrupting the BBB), to maintain efficacy while reducing systemic toxicity. The off-label uses of the medication, as well as the risks, including intracranial hemorrhage, were reviewed in detail. Informed written consent was provided by the parents, and the patient gave her assent. We performed intraarterial infusion of 2.5 mg/kg bevacizumab after hyperosmotic BBB breakdown. There were no acute complications from the procedure.

**Postintervention Course**

Within 12 hours of intraarterial bevacizumab administration, the patient experienced complete resolution of her previously intractable headaches (now a 0 on the 10-point scale). She also felt subjectively stronger on her right side. She was started on a long steroid taper under supervision of the Endocrine Service.

Two months later, brain MRI revealed an 82% decrease in FLAIR signal and a 6% decrease in contrast enhancement (Fig. 1 left and right, respectively). There was an associated markedly reduced mass effect. MRI of the brain at 5 months revealed 78% decreased FLAIR signal and 22% decreased contrast enhancement (Fig. 1 left and right, respectively).

At 7 months, the patient exhibited a progressive and prominent improvement in her proximal right arm and leg strength. She was able to walk long distances with the help of an electronic drop-foot unit. She has continued to lose weight (13 pounds so far). She experiences occasional moderate headaches (5–6 on a 10-point scale) that would last several hours, but the headaches are not the constant.
very severe ones she was experiencing before treatment. Her emotional lability resolved and she was able to go back to school.

Case 2

History and Examination

An 11-year-old right-handed girl presented with right hemiparesis, headache, nausea, and vomiting. Head CT scanning revealed intraparenchymal and intraventricular hemorrhage and hydrocephalus. She required temporary external ventricular drainage. MRI imaging demonstrated a Spetzler-Martin Grade II AVM of the corpus callosum. She underwent embolization 2 months later with only partial obliteration. She then underwent SRS for the residual AVM nidus with an 18-Gy radiation dose prescribed to a volume of 4.3 cm³.

Within the 1st month after treatment, the patient developed moderate headache, which responded to a 2-week course of steroids. Six months after SRS, she developed mild to moderate headaches, which initially responded to ibuprofen. MRI showed evidence of mild vasogenic edema surrounding the AVM site. Eight months after radiosurgery, she developed intractable headaches that were associated with nausea and vomiting. Repeat MRI demonstrated worsening cerebral edema with new contrast enhancement consistent with RN in the left frontal lobe (Fig. 2). There were no associated motor or sensory symptoms. A 21-day course of steroid treatment, starting with 24 mg per day of dexamethasone, was initiated. This steroid treatment could not, however, be tapered below 8 mg daily over the ensuing 3 months because of the patient’s recurrent severe headache, nausea, and vomiting. Courses of pentoxifylline and vitamin E were tried without success. Over time, she developed significant steroid-related symptoms. She also required a hospitalization for fluid overload. In less than 3 months, the patient gained over 30 pounds and had a body mass index of 25.5. As a result of her symptoms, she too had to withdraw from school.

Intervention

Based on the initial experience with Case 1, we discussed the option of intraarterial bevacizumab treatment. The off-label uses of the medication and the risks, including intracranial hemorrhage, were reviewed in detail. An informed written consent form was signed by the parents, and the patient provided assent. The patient underwent intraarterial infusion of 2.5 mg/kg bevacizumab into the left internal carotid artery after hyperosmotic BBB breakdown. There were no complications.

Postintervention Course

Immediately after intraarterial bevacizumab administration, the patient experienced complete resolution of her intractable headache, and within 4 weeks she was successfully weaned off steroids. Brain MRI 3 months later revealed a 74% decrease in FLAIR signal volume and a 33.6% decrease in contrast enhancement (Fig. 2 left and right, respectively). At 8-month clinical follow-up examination, the patient was free of headaches and was back in school.

Discussion

Stereotactic radiosurgery has become integral in treatment of brain tumors and AVMs. In up to 10% of cases, this can lead to RN with significant surrounding vasogenic edema and mass effect. Medical treatment for RN includes steroids, vitamin E, pentoxifylline, and hyperbaric oxygen. Up to 20% of cases are medically refractory and the patients experience progressive neurological decline and disabling headaches.
Role of VEGF in RN

Vascular endothelial growth factor is a tyrosine kinase that plays an important role in angiogenesis and modulation of vascular permeability. VEGF-A binds with high specificity to VEGF receptor–1 and VEGF receptor–2 on vascular endothelial cells. These modulate downstream signaling pathways affecting various cellular processes. VEGF has recently been implicated in the pathophysiology of RN. Reactive astrocytes immediately surrounding the necrotic core in RN are strongly positive for VEGF by immunohistochemistry. It has been postulated that radiation causes microvascular injury leading to hypoxia. Hypoxia-induced VEGF upregulation then drives an increase in vascular permeability, leading to the extensive vasogenic edema seen in RN.

Bevacizumab Therapy for RN

Bevacizumab (Avastin, Genentech BioOncology) is a recombinant humanized version of a murine anti–human VEGF monoclonal antibody. It binds circulating VEGF receptors with high specificity, blocking the downstream signaling cascade. Bevacizumab was originally developed and tested as an antiangiogenic treatment for various solid tumors.

More recently, intravenously administered bevacizumab was shown in a blinded, placebo-controlled, randomized trial (n = 14) to be effective in the treatment of refractory RN after radiation therapy in brain tumors. Patients received 7.5 mg/kg bevacizumab intravenously every 3 weeks for 4 cycles. All patients receiving bevacizumab, and none of the patients receiving placebo, had significant clinical and radiographic improvement. This improvement was durable, persisting at 10 months in 9 (92%) of 11 patients. There was, however, a high rate of adverse events (55%), and major adverse events occurred in 27%. There have since been a handful of AVM RN cases in adults successfully treated with intravenous bevacizumab.

There are significant known side effects of bevacizumab including gastrointestinal tract perforation, deep venous thrombosis, venous sinus thrombosis, pulmonary embolus, intracranial hemorrhage, wound dehiscence, and severe hypertension. These complications are common in the antiangiogenic class of drugs and reflect systemic exposure to bevacizumab. Current intravenous bevacizumab regimens use a dose of 7.5 mg/kg every 3 weeks for 4 cycles. Using a lower bevacizumab dose may lower systemic exposure and decrease complication rates, providing a rationale for further study. To maintain therapeutic efficacy of the drug at the lower dose, however, its ability to penetrate the brain must be improved. There are two ways to increase drug penetration into the brain: intraarterial route of administration and blood-brain barrier disruption.

Intraarterial Delivery

Intraarterial therapy decreases volume dilution of the drug in the circulation and reduces first-pass degradation through proteolytic catabolism. Superselective intraarterial injection of 99mTc–hexamethylpropyleneamine oxime (Ceretec) into human cerebral arteries achieves a concentration of radiotracer in brain tissue 50 times higher than with intravenous injection. In clinical studies of cerebral chemotherapy, the concentration delivered to the tumor via intraarterial injection versus intravenous injection of chemotherapeutic agents was 5 times higher with hydro soluble drugs and up to 50 times higher with liposoluble drugs. Intraarterial therapy does, however, involve an invasive angiographic procedure. Risk of significant procedure-related morbidity is less than 1%, but can include stroke, allergic reaction to contrast dye, kidney impairment, and bleeding from the access site.

Blood-Brain Barrier Disruption

The BBB is a selective permeability barrier that blocks entry of many drugs into the brain. Bevacizumab is a monoclonal antibody with a high molecular weight (149 kD). There is convincing evidence in the literature that the concentration in the brain of high–molecular weight molecules can be significantly increased after osmotic BBBD. Several tumor clinical trials have shown that localization of monoclonal antibodies to the brain is poor.
Intraarterial Bevacizumab Infusion After Osmotic BBBD for Treatment of RN

We used a combination of an intraarterial route of drug delivery and BBBD to reduce the bevacizumab dose while maintaining efficacy. This is supported by the durable clinical and radiographic response in our 2 patients after a single 2.5-mg/kg dose of bevacizumab. This approach may reduce the incidence of serious systemic toxicities compared with the intravenous bevacizumab regimen (7.5 mg/kg every 3 weeks for 4 cycles).

The safety of intraarterial bevacizumab treatment after hyperosmotic BBBD was recently established in a small series of patients with malignant glioma. This was done through superselective injection of intracranial tumor arterial pedicles for the purpose of antitumor effects. Dose escalation was performed from 2 mg/kg to 15 mg/kg without reaching the maximal tolerated dose. There was a significant decrease in the contrast-enhancing and FLAIR signal characteristics of the tumor and surrounding brain at 1 month after treatment. Overall toxicity in this cohort was comparable to that in previous reports for intravenous bevacizumab therapy. In addition, hyperosmotic BBB breakdown followed by intraarterial bevacizumab administration did not cause any direct neurotoxicity; there were no cases of intracranial hemorrhage. In a different study, the same authors also performed extensive cost analysis of bevacizumab therapy for recurrent glioblastoma. They showed a 58% total cost savings for bevacizumab treatment when it was administered intraarterially as compared with intravenously. This savings was achieved despite the fact that the authors used a 15-mg/kg intraarterial dose of bevacizumab, which is 6 times higher than the concentration used in the present case series.

Herein, we have described 2 pediatric patients with highly symptomatic RN in the brain after SRS for the treatment of cerebral AVMs. The RN was refractory to all accepted medical therapies. Both patients were steroid dependent for a prolonged period and had developed severe cushingoid features. Both had suffered a significant decline in quality of life, suffering severe headaches and needing to withdraw from school. In both instances, the patients made a remarkable clinical and radiographic improvement after receiving a single low dose of intraarterial bevacizumab following hyperosmotic BBB breakdown. To our knowledge, this report marks the first demonstration of the successful treatment of cerebral RN using targeted intraarterial bevacizumab therapy.

Conclusions

We propose earlier consideration of bevacizumab treatment to avoid neurological morbidity and severe adverse effects of long-term steroid treatment. We have shown that intraarterial bevacizumab after hyperosmotic BBB breakdown appears to be safe, effective, and durable treatment for medically refractory RN in the brain. Advantages of the intraarterial route over the intravenous route may include higher concentration of drug delivery to the targeted brain tissue, decreased systemic toxicity because of the much smaller dose of bevacizumab used, and the significantly lower cost. The main limitation of the present report is the very small number of patients. Future prospective clinical trials comparing intravenous and intraarterial bevacizumab treatment in this setting deserve consideration.

References

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