Noninvoluting congenital hemangiomas (NICHs) are described as part of the spectrum of vascular tumors. They differ from rapidly involuting congenital hemangiomas, which proliferate rapidly in the first year of life and then gradually regress over the following 5 to 10 years. NICHs in contrast, fail to involute and histologically stain negative for glucose transporter protein 1 (GLUT-1). Hemangiomas are proliferative vascular lesions characterized by increased endothelial cell turnover, and NICHs therefore can result in large and highly vascularized lesions whereby surgical intervention poses considerable hemorrhage risk. In such cases, presurgical embolization can play a significant role. Pre-embolization work-up includes careful evaluation of the lesion using catheter angiography to assess for feeding and draining vessels. We present a case of a 3-year-old boy with a giant vertex hemangioma who developed systemic complications following embolization with Onyx (ethylene vinyl alcohol copolymer) and Glubran (N-butyl-2-cyanoacrylate).

Case Report

This 3-year-old boy was born with a small vertex scalp swelling. MRI performed shortly after birth showed that the scalp swelling was a highly vascular lesion that was completely extracranial. There was no particular abnormality of the brain or spine. The lesion grew steadily over the course of the first year and biopsy was performed when the patient was 1 year old. The biopsy showed fibroconnective tissue containing irregular, enlarged and dilated thin-walled vessels lined by plump endothelial cells. There was no evidence of atypia or malignancy. Immunohistochemical staining was negative for GLUT-1 and podoplanin, suggesting a NICH (Fig. 1).
Possible toxicity after embolization of hemangioma

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Sultant chronic hypochromic microcytic anemia, a trial of propranolol was commenced. Prior to propranolol treatment the lesion measured approximately 15 × 15 cm. There was no reduction in size after 6 months of treatment at a peak dose of 2.5 mg/kg/day, and elective resection of the hemangioma under the combined management of a neurosurgeon and craniofacial surgeon was therefore scheduled. This procedure was abandoned due to significant intraoperative hemorrhage, and subsequently a presurgical embolization was planned. Four-vessel cerebral angiography showed no evidence of cerebral vascular abnormality or vessels communicating with or feeding the scalp hemangioma (Fig. 2A). The patient subsequently underwent a course of percutaneous intralesional embolization therapy (Fig. 2B) over 3 sessions. In the first session 8 ml of ethylene-vinyl alcohol copolymer (Onyx) was injected, resulting in embolization of approximately one-quarter of the lesion. In the second session 4 ml of Glubran was mixed in 30 ml lipiodol, a radiopaque agent, and injected into the lesion. In the third session 17 ml of Glubran mixed in 60 ml lipiodol was injected (Fig. 2C and D). Five days following the second session the patient complained of nonspecific leg pains, which resolved approximately 5 days later without any specific intervention.

A decision was made to electively resect the lesion 2 weeks following the final embolization, but 5 days following the third embolization session, he developed sudden-onset ataxia, tremor, and lethargy. The tremor was symmetrical and action induced, involving head, trunk, and limbs. There was no nystagmus or dysarthria. The patient’s muscle strength and deep tendon reflexes were normal. He was irritable but his cognition was intact.

A CT scan of the brain showed subtle bilateral hyperdensities in the caudate nuclei, thalami, and cerebellar cortex (Fig. 3D). MRI of the brain was considered, but as there was concern regarding consequences of an electromagnetic heating effect of the intralesional Glubran, this was not undertaken. A chest radiograph showed diffuse interstitial changes consistent with interstitial pneumonitis (Fig. 3A). A CT scan of the thorax showed similar diffuse interstitial changes bilaterally with multiple areas of density in the perihilar and subcarinal regions (Fig. 3B).

Despite this finding, there were no respiratory signs or symptoms. A CT scan of the abdomen did not show any remarkable features (Fig. 3C).

Based on the clinical presentation and radiological findings, the possibility of toxicity secondary to Onyx or Glubran, or an immune-mediated reaction to either was considered, with an embolic phenomenon less likely due to the delay in onset of symptoms. Urgent resection of the hemangioma was scheduled.

Intraoperatively, the patient was positioned prone, and the hemangioma was carefully resected off the occiput and the underlying skull defect. No dural attachment was noted during resection and the skin was closed directly (Fig. 4).

On the 1st postoperative day, neurological symptoms worsened. Tremor was more pronounced but was only action induced. The patient was irritable and unable to stand or feed himself but had normal strength and preserved cognition.

Following a stat dose of intravenous dexamethasone (4 mg), the symptoms decreased in severity. Dexamethasone was continued in addition to a 5-day course of 2 g/kg intravenous immunoglobulin (IVIG). Clonazepam was prescribed for 72 hours for agitation and tremor. The patient’s symptoms resolved over the following days and he was discharged home on a tapering dose of oral prednisolone. There was no blood eosinophilia, and histological examination of the resected hemangioma showed no eosinophilia or vasculitis. In retrospect, it would have been valuable to have had immunological investigations, but due to the sudden presentation, empirical treatment had to proceed, and fortunately the patient responded rapidly.

Neurological examination, a chest radiograph, and an
MRI study of the brain performed 6 months later showed no abnormality.

**Discussion**

This report describes an unexpected complication of Onyx and Glubran treatment of hemangioma. These materials are the most widely used liquid embolic agents in the world, but their adhesive properties and liquid nature require special precautions. Clinical manifestations in our patient may have been immune mediated. A prior report by Quinn et al. highlighted the potential for Glubran to produce an eosinophilic vasculitis in embolized intracranial arteriovenous malformations (AVMs). Additionally, other portions of nonembolized AVMs showed similar eosinophilic infiltrate, arguing that this is probably a nonfocal reaction. While none of these patients showed signs of systemic vasculitis, the authors concluded that long-term exposure to Glubran and subsequent development of an eosinophilic vasculitis in these patients may lead to further neurological sequelae or an increased risk of vessel rupture. Furthermore, a humoral immune response against Glubran may contribute to clinical symptoms.

We hypothesize that the neurological symptoms and radiological findings could be explained by an immune-mediated process and therefore the patient was treated with steroids and IVIG. One week following discharge, symptoms had resolved and the child was neurologically normal. Although there were no blood or histological markers of an immune-mediated or allergic process, such a mechanism cannot be excluded.

While risks such as embolization to the venous system, intracranial hemorrhage, and pulmonary embolism have been reported in Glubran embolization of brain AVMs, an embolic phenomenon was not considered likely in this case due to the delay in onset of symptoms. Embolic events are more often instantaneous. Furthermore, there were no direct communicating or feeding vessels between the hemangioma and the systemic circulation. As the procedure was performed under continuous fluoroscopy, any potential migration of the Onyx or Glubran from the embolized area should have been visible. Relatively invisible communications between the hemangioma and systemic circulation may have facilitated the systemic toxicity or immune-mediated effect rather than the spread for emboli to multiple areas.

Despite the complications mentioned in the literature to date, none relate to the treatment of an extracranial lesion, as reported here. A definitive safe maximum dose has not yet been reported and it is largely driven by subsequent radiological appearance of the lesion as well as by the experience of the intervention radiologist. Murai et al. reported successful outcomes in 10 patients who underwent embolization for cerebellar hemangiomas with no reported side effects. The specific dose used is not stated. Effective therapy of Glubran treatment in laryngopharyngeal hemangioma has also been reported with a dose of 0.5–2 ml.

In our case the dose used allowed successful and uncomplicated resection of what was initially a highly vascular lesion.

In summary, we present a significant and previously unreported side effect of Onyx and Glubran manifesting as a severe tremor, possibly immune mediated, following treatment of a giant congenital hemangioma. This complication should be borne in mind when using these embolic agents.

**References**

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Disclosures
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