Intracranial intraaxial cerebral tufted angioma: case report

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Tufted angioma (TA) is a rare, slow-growing, vascular lesion that commonly presents as a solitary macule, papule, or nodule arising in the soft tissues of the torso, extremities, and head and neck in children and young adults. Adult-onset cases have been infrequently reported. While typically benign, TAs may be locally aggressive. Complete physical examination and hematological workup are recommended in patients with TA to exclude the presence of Kasabach-Merritt phenomenon (KMP). The authors describe the case of a 69-year-old man with a contrast-enhancing frontal lobe lesion, with surrounding vasogenic edema, which was treated by gross-total resection. Characteristic histological features of a TA were demonstrated, with multiple cannonball-like tufts of densely packed capillaries emanating from intraparenchymal vessels in cerebral cortex and adjacent white matter. Tumor recurrence was detected after 4 months and treated with adjuvant Gamma Knife radiosurgery. To the extent of the authors’ knowledge, this case illustrates the first report of TA presenting in an adult as an intracranial intraaxial tumor without associated KMP. The fairly rapid regrowth of this tumor, requiring adjuvant treatment after resection, is consistent with a potential for locally aggressive growth in a TA occurring in the brain.

https://thejns.org/doi/abs/10.3171/2016.10.JNS162207

KEY WORDS tufted angioma; brain tumor; intracranial; Kasabach-Merritt phenomenon; kaposiform hemangioendothelioma; vascular disorders

Tufted angioma (TA) is a rare, slow-growing, vascular tumor derived from capillary and lymphatic endothelium. Originally, TA was termed an angio-blastoma17 or progressive capillary hemangioma.14 In 1976, Jones coined the name “tufted angioma” based on its histological features of multiple, discrete, cannonball-like lobules (“tufts”) of densely packed capillaries surrounded by crescentic spaces.6,12 TA usually presents as a solitary, expanding, erythematous, or violaceous soft-tissue macule, papule, or plaque.19 Generally, TA is recognized as a benign lesion that may nonetheless be locally aggressive but without reported evidence of distant metastases. However, multifocal cases have been reported.20 Children and young adults are predominantly affected by TA, with adult onset described infrequently.9,50

To date, the etiology and pathogenesis of TA remain uncertain, although some believe TA and kaposiform hemangioendothelioma (KHE) are part of a continuum of the same pathological process.9,19 Interestingly, both TA and KHE may be associated with Kasabach-Merritt phenomenon (KMP), a clinical constellation of thrombocytopenia, consumptive coagulopathy, and purpura associated with larger lesions (> 5 cm).8,9,19,21

Together, TA/KHE have been known to arise in a number of anatomical locations including the soft tissues of the extremities, the torso, mediastinum, retroperitoneum, head and neck region, and rarely oral mucosa.5,8 Cases of KHE involving the internal/external auditory canals and the tentorium cerebelli, without evidence of parenchymal involvement, have been reported.2,3,10 To our knowledge, no reports exist of TA/KHE arising within brain parenchyma. In this paper we describe the case of a 69-year-old man with a contrast-enhancing frontal lobe lesion, with surrounding vasogenic edema, which was treated by gross-total resection. Characteristic histological features of a TA were demonstrated, with multiple cannonball-like tufts of densely packed capillaries emanating from intraparenchymal vessels in cerebral cortex and adjacent white matter. Tumor recurrence was detected after 4 months and treated with adjuvant Gamma Knife radiosurgery. To the extent of the authors’ knowledge, this case illustrates the first report of TA presenting in an adult as an intracranial intraaxial tumor without associated KMP. The fairly rapid regrowth of this tumor, requiring adjuvant treatment after resection, is consistent with a potential for locally aggressive growth in a TA occurring in the brain.

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old man with a medial frontal lobe intraaxial TA without associated KMP, treated using gross-total resection and subsequent adjuvant Gamma Knife radiosurgery (GKRS).

**Case Report**

**History and Examination**

A 69-year-old man presented with mild subjective memory complaints. There was no family history of hereditary disease or malignancy. No laboratory abnormality was found, including any platelet abnormality. Precontrast CT of the head demonstrated a hypodense lesion in the right superior medial frontal lobe with no evidence of hemorrhage or calcification (Fig. 1A). Postcontrast CT demonstrated avid curvilinear enhancement located medially (Fig. 1B). There was no bone involvement. Brain MRI was notable for a right frontal, parasagittal, avidly enhancing lesion lying flat along the falx cerebri and involving the cortex and subcortical white matter (Fig. 1C and D). The enhancement pattern was noted to be curvilinear and ribbon-like, outlining the cortex. A hyperintense T2 FLAIR signal was noted surrounding the enhancing lesion and confined to the right superior frontal lobe with mild associated gyral thickening, suggesting the presence of surrounding vasogenic edema (Fig. 1E). There was no associated diffusion restriction or susceptibility. The initial differential diagnosis included: dural-based lesions such as meningioma, given its proximity to the falx; brain metastasis, although the patient had no evidence of systemic cancer; cortically based enhancing lesions such as encephalitis, inflammation, subacute infarct, and seizures; and low-grade neoplasms such as dysembryoplastic neuroepithelial tumor, ganglioglioma, pilocytic astrocytoma, and pleomorphic xanthoastrocytoma.

**Operation**

Repeat imaging after 3 months revealed that while the amount of edema was similar in comparison with the prior scan, the lesion had enlarged by 2.5 mm in the axial plane and 1.0 mm in the coronal plane (Fig. 2). Surgery was performed using a right parasagittal craniotomy with frameless stereotactic guidance. Dissection was carried down along the interhemispheric surface of the right medial frontal lobe until obvious pathological tissue was visually identified. Of note, there was no dural attachment of the tumor. Intraoperative ultrasound was used to confirm the location of the lesion, combined with frameless stereotactic navigation. Subsequently, the lesion was entered and debulked. An intraoperative frozen-section diagnosis of angiocentric neoplasm was rendered (data not shown). The mass was circumferentially dissected and removed except for a small portion anteriorly attached to a prominent artery and vein sulcal complex. These vessels were
preserved, and microdissection techniques were applied until there was no identifiable tumor remaining. However, residual microscopic disease attached to these vessels was likely.

Histological Analysis

Sections of the lesion stained with H & E showed a vascular neoplasm of moderate cellularity growing in small, tightly packed, rounded nests (“tufts”) of angiomatous proliferations along parenchymal vessels in cerebral cortex and the adjacent white matter (Fig. 3A and B). Each nest contained clusters of plump endothelial cells surrounding small, slit-like capillary lumina (Fig. 3C). Cells surrounding the vascular channels had oval- to spindle-shaped nuclei with fine chromatin and only mild nuclear pleomorphism, and lacked epithelioid features. Apoptotic cells and rare small foci of necrosis were noted (Fig. 3C). In some foci, vessel walls were partially or completely sclerotic (Fig. 3D). The endothelial cells showed immunoreactivity for several markers, including CD31, CD34, FLI1 (Fig. 4A–C), and focally for GLUT1 (likely representing feeding vessels, data not shown). Cells in the nests were otherwise faintly positive for progesterone receptor, and negative for STAT6, inhibin A, pan-cytokeratin, epithelial membrane antigen, and D2–40 (data not shown). Many of the cells interspersed between vascular channels appeared to be pericytes and labeled with anti–smooth muscle actin (Fig. 4D). Scattered mitotic figures were noted, and Ki 67 labeling ranged from 9.5% to 25.6% of cells (Fig. 4E). In addition, a mixed inflammatory infiltrate was noted, including CD68-positive histiocytes and CD45-positive lymphocytes observed between neoplastic cells within the nests (data not shown). The adjacent brain parenchyma showed marked reactive gliosis, including astrocytes highlighted by an immunostain for glial fibrillary acidic protein (GFAP; Fig. 4F) and CD68-positive microglia (data not shown).

Postoperative Course

Postcraniotomy, the patient was maintained on a 1-week steroid taper. Postresection MRI demonstrated gross-total resection with no residual contrast enhancement in the surgical bed (Fig. 5) and mildly reduced T2 FLAIR signal within the surrounding parenchyma. As the potential biological behavior of an intracranial intraaxial TA is unclear, routine surveillance MRI in 3–4 months was recom-
Intracranial tufted angioma

In this paper we describe, to the best of our knowledge, the first case of an intracranial intraaxial TA. The differential diagnosis for TA includes KHE, capillary hemangioma, infantile hemangioma, glomeruloid hemangioma, vascular malformations, Kaposi sarcoma, hemangioblastoma, and hemangiopericytoma. Rare reports do exist of KHE arising from the temporal bone and the tentorium cerebelli, and these typically show locally invasive growth. TA is generally regarded as closely related to KHE, named for its histological resemblance to Kaposi sarcoma and also containing infiltrating nodules with slit-like or crescentic vessels lined by spindled endothelial cells. Some believe that TA and KHE are part of a continuum of the same pathological process. The lesions of KHE appear very similar histologically to those of TA, but are more infiltrative with larger coalescing nodules. In addition, the lesions of KHE also contain epithelioid cells. In both lesions, platelet-rich fibrin thrombi may accumulate, leading to KMP. However, KMP develops in more than 70% of patients with KHE and in only 10% of patients with TA.

In the present case, there were no associated laboratory abnormalities concerning for KMP. Capillary hemangiomas are also common cutaneous lesions, but they have only rarely been reported in the brain, where they predominantly occur in an extracranial location with circumscribed growths of capillaries surrounded by fibrous septae. In a systematic review of the literature, Mirza et al. found only 3 intracerebral cases of capillary hemangiomas among 14 adult patients. However, none of these intraaxial capillary hemangiomas exhibited a lobular architecture; rather, they had a papillary pattern and areas with capillaries dispersed in an edematous collagenous stroma. Other variants of hemangiomas in the differential diagnosis include infantile hemangiomas and glomeruloid hemangiomas. Infantile hemangiomas consist of closely packed capillary spaces with plump endothelial cells that express GLUT1; however, they do not display a cannonball-like architecture as in the present case. Glomeruloid hemangiomas, which are aggregates of vessels that resemble renal glomeruli, are usually associated with POEMS syndrome (polyneuropathy, organomegaly, endocrinopathy, monoclonal gammopathy, and skin changes) and Castleman disease. Kaposi sarcoma lesions show jagged, infiltrative, interconnected vascular channels with interspersed plasma cells. A diagnosis of hemangioblastoma or hemangiopericytoma was excluded by negative immunostains for inhibin A and STAT6, respectively. Taken together, the morphological and immunophenotypic features of this lesion are most consistent with TA.

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The lesion in the present case had mitotic figures and an elevated Ki 67 index, indicative of a proliferative process. This supports the evidence of tumor progression witnessed on serial MRI. Of note, an elevated proliferation index is not unusual in TA or KHE, and does not by itself lead to a diagnosis of angiosarcoma. In the current case, the tumor’s discrete growth pattern and lack of marked cellular atypia do not support a diagnosis of angiosarcoma. While
microscopic foci of residual TA after surgery may have allowed for recurrence in the absence of adjuvant therapy, the fairly rapid regrowth of this lesion within 4 months is consistent with the known locally aggressive growth potential for TA.

TA is a rarely described entity, and treatment modalities have been determined without randomized controlled trials or prospective studies. Single case reports and several small series form the foundation for recommendations, which include observation, surgery, medical therapies, arterial embolization, pneumatic compression, and radiation. Complete resection, however, remains the treatment of choice for small, localized lesions. Postoperative surveillance is recommended as recurrence of both TA and KHE has been reported. Malignant transformation of these lesions has never been reported. No evidence currently exists regarding the efficacy of GKRS for these lesions.

Conclusions

We have presented a case of TA, with no associated laboratory abnormalities concerning for KMP, arising within the parenchyma of the right frontal lobe. The tumor rapidly recurred after radiographically confirmed complete resection requiring adjuvant GKRS. This is the first reported case of this type of tumor in an intracranial, intraxial location. In addition, this report provides a novel description of the morphological growth pattern and immunophenotypic features of an unusual intraxial angioma. TA should be considered in the differential diagnosis of intracranial, intraxial vascular tumors and resection should be considered the treatment of choice, particularly

FIG. 7. GKRS plan. A single treatment plan was devised to encompass the area of recurrence along the axial (A), coronal (B), and sagittal (C) planes inferior to the margin of the old resection bed, as well as incorporating the entirety of the old surgical bed and dural margin to minimize the risk of tumor progression. Target volume was 607.8 mm³. Given the irregular shape of the lesion tracking along the interhemispheric fissure and abutting the resection cavity, a total of 12 4-mm shots with sector blocking were outlined to deliver 18 Gy to the 50% isodose line. A 3D rendering of the skull is included in the plan, with red representing the assumed shape of the skull for dosimetry purposes (D). The rectangle in panel A is the “shot view” and shows the 13th shot is not set, i.e., there are only 12 shots set. Figure is available in color online only.

FIG. 8. Follow-up imaging 4 months after GKRS. Postcontrast axial (left) and coronal (right) MR images demonstrate stable enhancing disease at the posterior inferior tumor bed.
in the absence of KMP. GKRS may be considered for recurrence.

References


15. Mentzel T, Wollina U, Castelli E, Kutzner H: [Tufted hemangioma. Clinicopathologic and immunohistologic analysis of 5 cases of a distinct entity within the spectrum of capillary hemangioma.] Hautarzt 47:369–375, 1996 (Ger)


Disclosures

Dr. Wang has served as a consultant to Abbvie, Merck, and Doximity.

Author Contributions


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