Malignant ossifying fibromyxoid tumor of the calvaria: illustrative case

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BACKGROUND Ossifying fibromyxoid tumor (OFMT) is a rare entity of soft tissue tumor that most commonly occurs in the subcutaneous tissues of trunk or extremities with occasional cases involving the head and neck; however, primary involvement of the skull has not been reported. While historically considered slow-growing benign to intermediate malignant, few cases of atypical or malignant features have been described.

OBSERVATIONS Herein, the authors present a case of malignant OFMT with primary skull and transcranial extension. The tumor caused lytic calvarial destruction with intra- and extracranial soft tissue components. Gross total resection was performed, and histopathology revealed malignant OFMT with 40 mitoses per 50 high-power fields and moderate nuclear atypia.

LESSONS OFMT can rarely occur in the head and neck and, as reported herein, may involve the skull with intracranial extension. While no uniformly recognized histological criteria for malignancy exist, a three-tiered classification has been proposed: typical, atypical, and malignant, based on features such as hypercellularity, mitotic activity, infiltrative growth, and/or nuclear atypia. Malignant variants should be considered along the high-grade sarcoma spectrum with elevated risk for recurrence or metastatic spread. Routine adjuvant radiotherapy is not typically recommended; however, surveillance imaging is advised.

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KEYWORDS malignant ossifying fibromyxoid tumor; sarcoma; calvaria; skull; oncology; ossifying fibromyxoid tumor

Ossifying fibromyxoid tumor (OFMT) is a rare and distinct mesenchymal soft tissue tumor entity of unknown histogenesis. It was first described in 1989 by Enzinger et al. as “ossifying fibromyxoid tumor of soft parts.” While historically considered benign or indeterminate/borderline malignant potential, frank malignant variants have been recognized and are considered part of the high-grade sarcoma spectrum. While the exact differentiation cell line remains unknown, there is some evidence of putative Schwannian/neuroectodermal etiopathogenesis or myoepithelial/cartilaginous differentiation, with a “scrambled phenotype” being a leading hypothesis.

Most frequent affected regions by this tumor subtype include the subcutaneous soft tissues of the trunk and extremities, with proximal more common than distal locations. Occasional cases involving the head and neck region have been described, affecting mainly the anterior and posterior neck soft tissues, face, and oral cavity (e.g., cheek, lip, chin, soft palate, nose, zygoma/parotid, submental and submandibular region, and tongue). Rarely, involvement of the scalp has been described. However, to our knowledge, this is the first reported case with primary calvarial bone involvement and transcranial extension.

Illustrative Case

The patient is a 25-year-old female with a medical history of polycystic ovarian syndrome and hypothyroidism who presented with gradually enlarging left scalp swelling over a period of 12 months. On exam, there was scalp expansion over the left temporoparietal region; the overlying skin was unremarkable without any overt changes. Prior to neurosurgical referral, the following workup had been performed: An ultrasound examination showed a cystic scalp mass with low-level internal echoes, foci of hyperechogenicity, internal vascularity, and thin hypervascular rim. A punch biopsy to the depth of the superficial subcutaneous fat showed only sparse perivascular lymphocytic inflammation and slight sclerosis of dermal collagen but was otherwise unremarkable. A plain skull radiograph was obtained that showed a 3-cm radiolucent lesion at the left parietal calvaria. Magnetic resonance imaging (MRI) of the brain

ABBREVIATIONS CT = computed tomography; HPF = high-power field; MRI = magnetic resonance imaging; OFMT = ossifying fibromyxoid tumor.

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without and with contrast was obtained, and the patient was referred for neurosurgical evaluation. Neurological exam was unremarkable. The MRI revealed a lytic transcranial left parietal mass lesion with intra- and extracranial soft tissue extension and lytic destruction of the calvarial skull in the area. Dimensions of the mass were 5 cm anteroposterior by 3.8 cm transverse by 5.6 cm craniocaudal. The intracranial component appeared limited by the dura. The cortical surface was depressed by approximately 1 cm without evidence of brain invasion or adjacent perifocal edema. The mass showed homogenous intermediate signal intensity on T1- and T2-weighted imaging, no diffusion restriction, and uniform avid contrast enhancement (Fig. 1).

The patient underwent an elective left parietal craniectomy and methylmethacrylate cranioplasty (Fig. 2). A linear incision was planned to span the lesion extending from normal inferior to normal superior bony margins; the incision was opened with close attention not to violate the capsule of the mass. The periosteum was carefully freed up and the temporalis muscle was partially reflected. Two burr holes were made and a craniectomy was performed circumferential to the lesion with a clear margin. The tumor was noted to be adherent to the underlying dura. A plane was carefully developed using blunt and sharp dissection. Gross total resection was performed en bloc, and the mass was sent for frozen and permanent pathology. After the frozen section revealed a potentially sarcomatous lesion, the underlying dura in contact with the lesion was excised with a margin and sent for permanent pathology. A dural allograft patch was then secured in a standard watertight fashion. A methylmethacrylate cranioplasty implant was shaped to size and form and was secured in place using standard cranial plating technique. Incision was closed in the standard fashion.

The outer surface of the skull was smooth with an attached irregularly shaped soft tissue mass measuring 6.8 × 6.5 × 2.7 cm. The mass showed dense fibroconnective tissue, grossly encapsulated by a fibrous capsule with thin rim of ossification at the edge of the tumor. The inner surface of the skull was concave and cracked with a central irregularly shaped defect, measuring 3.8 × 3.1 cm, with protruding soft tissue lesion in continuity with the external component of the mass (Fig. 3). The inked margins of the resected mass were negative for tumor. The cut surface of the tumor was tan-white, fleshy, and glistening with scant focal areas of hemorrhage. The surrounding normal appearing bone was unremarkable; there was no evidence of satellite lesions. The resected dura showed microscopic foci of adherent tumor without evidence of transgression or invasion. Histopathological analysis (Fig. 4) revealed a lobulated spindle cell neoplasm in chondromyxoid matrix with a characteristic ossifying rim. There was no significant necrosis. However, moderate nuclear pleomorphism and atypia were apparent, and there were scattered giant cells and numerous mitotic figures (40 per 50 high-power fields [HPFs]), consistent with a diagnosis of malignant OFMT. Immunohistochemical stains were negative for S100, CD34, Desmin, Actin smooth muscle, and CKAE1/AE3. Staining was also negative for STAT6/MUC4. INI1 showed retained nuclear expression throughout the lesional tissue. There was patchy weak staining for CD99.

The postoperative course was unremarkable; the patient was monitored in the neuroscience intensive care unit overnight, transferred to the general neurosurgical ward on postoperative day 1, and discharged home on the morning of postoperative day 2. A

![FIG. 1. Preoperative imaging showing 5 cm by 3.8 cm by 5.6 cm transcranial left parietal mass with intra- and extracranial extension and central lytic destruction of the calvarial bone with approximately 1-cm dural/cortical displacement. A: Axial T1-weighted without contrast. B: Axial T1-weighted with contrast. C: Axial T2-weighted. D: Axial fluid-attenuated inversion recovery. E: Axial diffusion weighted imaging. F: Axial apparent diffusion coefficient. G: Coronal T1-weighted with contrast. H: Lateral skull radiograph showing a 3-cm radiolucent lesion at the left parietal calvaria.]
Discussion

Observations

The tumor presented herein showed morphologically classical appearance with dense fibroconnective tissue encapsulated by a fibrous capsule with an ossified rim. There was central destruction of the skull. Measuring 5 cm by 3.8 by 5.6 cm, the mass was adherent to the underlying dura causing approximately 1 cm of cortical/dural depression without any macro- or microscopic evidence of dural or brain invasion. Immunohistochemistry was negative for S100, CD34, desmin, smooth muscle actin, cytokeratin, STAT6/MUC4; INI1 was retained. The tumor presented in this report showed 40 mitoses per 50 HPFs, which is along the upper spectrum reported in the literature. The tumor also showed a moderate degree of nuclear atypia, but no necrosis was observed.

Lessons

OFMTs affect predominantly young to middle-aged adults with a slight male predilection; however, they can occur at any age, and a wide range has been documented (3 weeks to 83 years). It most commonly presents as a gradually enlarging circumscribed soft tissue mass. While typically painless, occasional reports of pain and/or paresthesias have been described. The clinical course is commonly described as a slowly enlarging lesion often over multiple
years (range 1–20 years), and lesions have been reported at varying sizes, ranging from 1 to 21 cm at presentation. Local recurrence rates have been reported in up to 22% of patients across the histological spectrum, with a typical interval around 10 years from index resection. Frequency of metastatic spread has been estimated at 5% of cases and more common in cases of malignant variants. Imaging appearance on MRI can be variable, including other soft tissue sarcomas such as low-grade fibroxoid sarcoma, sclerosing epithelioid fibrosarcoma, synovial sarcoma, Ewing sarcoma, osteosarcoma, chondrosarcoma, and malignant peripheral nerve sheath tumor but also ossifying epithelioid hemangioendothelioma or ossifying hematomas.

Histologically, OFMT is characterized by loose fibroxoid stroma with dense lobules of relatively uniform polygonal spindle cells in nests, cords, or trabeculae with fibrous septations. The cells are characterized by distinct cell membranes, variably scant eosinophilic cytoplasm, and round to oval vesicular nuclei with conspicuous nucleoli. OFMT is frequently positive for S100 and is more commonly expressed in typical than atypical or malignant variants (88% versus 75% versus 42%, respectively, in one study). Similarly, desmin is commonly expressed, staining positive in approximately half of typical/atypical cases and approximately one-quarter of malignant variants. Additionally, OFMT can variably express GFAP, vimentin, smooth muscle actin, cytokeratins, epithelial membrane antigen, neurofilament, CD56, or collagen II. Loss of INI-1 (commonly in mosaic pattern) or overexpression of EAAT4 and MUC4 are also not infrequent.

While there are no uniformly recognized histological criteria for malignancy, a three-tiered classification system has been proposed: typical, atypical, and malignant. Atypical features including hypercellularity, nuclear atypia, infiltrative growth pattern, and elevated mitotic activity have shown statistically significant correlations with clinically more aggressive courses, including elevated risk for local recurrence and, less commonly, distant metastatic spread. The presence of mitotic activity is highly variable, and in large series, has been reported as ranging from 0 to 40 per 50 HPFs. For classification as malignant OFMT, a cutoff of >12 mitoses per 50 HPFs has been proposed.

Focal necrosis, while not characteristic, can be observed in 10% to less than 20% of cases; vascular invasion occurs in approximately 10% of cases. Similar to other sarcoma entities, molecular associations and signature mutations continue to emerge: certain recurrent gene rearrangements involving the PHF1 gene have been reported, notably EP400-PHF1 translocation fusion genes. In a series of 41 OFMTs, Graham et al. detected PHF1 gene rearrangements in 49% of tumors, including 43% typical, 50% atypical, and 52% malignant variants. In a similar immunohistopathological cohort study of 39 OFMTs, Antonescu et al. identified additional recurrent gene fusions, increasing the yield to 85% of tumor samples (33 of 39 tested tumors). Additionally, INI-1/SMARCB1 deletions have been implicated in the pathogenesis of various sarcoma subtypes and have recently also been detected in a subset of OFMTs.

Herein, we present a rare case of malignant OFMT with primary skull involvement and transcranial extension, which, to our knowledge, has not previously been reported. No evidence of metastatic disease was apparent at the time of the index operation. Generally, treatment consists of gross total resection, favored en bloc, and complete excision is generally considered curative. Routine adjuvant radiotherapy is generally not recommended. Close follow-up is prudent, especially for atypical or malignant variants that should be regarded as sarcomas with potential for local recurrence on metastatic spread.
