Low-grade fibromyxoid sarcoma of the maxillary sinus
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Background. Low-grade fibromyxoid sarcoma (LGFMS) is a rare malignant tumor. Moreover, only 3 cases of LGFMS originating from the nasal cavity and/or paranasal sinuses have been published so far.

Methods. Two patients with histologically confirmed LGFMS of the maxillary sinus were primarily treated by open and endoscopic surgery. In one case adjuvant radiotherapy was indicated because of the uncertainty of the surgical margins.

Results. Both surgeries were technically demanding and accompanied by significant bleeding from the tumors. Despite the extensive interventions there were no postoperative complications and no significant morbidity of the patients. There is no evidence of the disease 148 and 65 months after the treatment.

Conclusions. Regardless of the excellent post-treatment results, based on our experience, preoperative embolisation of the tumor should be considered prior to the surgical resection.

Key words: sarcoma, low-grade fibromyxoid sarcoma, head and neck, maxillary sinus, endoscopic sinus surgery

INTRODUCTION

Low-grade fibromyxoid sarcoma (LGFMS) is a rare tumor counting less than 400 cases found in literature worldwide. The disease was described by Evans et al. in 1987, who published 2 cases of potentially metastasizing soft tissue tumor with deceptively benign histology, consisting of alternating myxoid and fibrous areas. Typical sites of the occurrence are deep soft tissues of the trunk and extremities. Only around 20 cases of LGFMS in the head and neck have be reported; however, most of them are part of case series studies without specification of the tumor location. To the best of our knowledge, there are only 2 brief reports describing LGFMS in the nasal cavity or paranasal sinuses and 1 case included in the patient cohort, but not discussed further. We present in detail 2 cases of LGFMS of the maxillary sinus that were diagnosed and treated at our department.

Case 1
A 57-year-old woman visited our outpatient department for 2 years persisting nasal obstruction on the left side. On the physical examination there was an uncertain finding of a hypertrophy of the mucous membrane of the middle concha and a tissue obturation of the upper part of the left choana. CT scan showed a pathological mass originating from the antrum of the left maxillary sinus and invading the nasal cavity (Fig. 1A). Based on the endoscopic examination and CT scan a malignant tumor was included in the differential diagnosis and the patient was indicated for endoscopy under general anesthesia. During the endoscopy, a greyish vasculated tumor on the lateral side of the nasal cavity was described. We evaluated that endoscopic resection was not possible in a radical fashion. A biopsy was taken but immediately after that severe bleeding from the tumor occurred and had to be controlled by an anterior nasal package. After the first procedure, MRI was performed during hospitalization, clarifying propagation of the tumor into the fossa pterygopalatina. The tumor was nonhomogeneous in signal density, but generally hyposignal (Fig. 1B,C). The acquired histology was described as chondroid chordoma. A decision on radical surgery was made, however, due to the bleeding experience and spreading of the tumor to the pterygopalatine fossa, an open resection through lateral rhinotomy with anterior and posterior ethmoidectomy, sphenoidectomy and resection of medial and posterior wall of the maxillary sinus was performed. Resection of the tumor from the pterygopalatine fossa, an open resection through lateral rhinotomy with anterior and posterior ethmoidectomy, sphenoidectomy and resection of medial and posterior wall of the maxillary sinus was performed. Resection of the tumor from the pterygopalatine fossa was accomplished in a piecemeal fashion. There was no postoperative complication. Unexpectedly, a final histological evaluation revealed low-grade fibromyxoid sarcoma. Due to the spread of the disease and uncertainty about surgical margins, adjuvant radiotherapy was indicated at a total dose of 60 Gy on the locality of the tumor. The patient was followed regularly using endoscopic examination and imaging methods. At the time of writing there has been no evidence of the disease 148 months after the surgery.

Case 2
A 74-year-old man was referred to our department with a suspicious mucocele of the right maxillary sinus. Nasal obstruction progressing over two months was the only symptom reported by the patient. During endoscopic ex-
Nevertheless, the bleeding was controlled endoscopically and the postoperative course was without any complication. A final histology report described the tumor as containing myxoid and cellular areas with star-shaped cells and rich vascularization. Following wide immunohistochemical testing the final diagnosis was LGFMS (Fig. 3). Despite the larger size of the tumor in comparison to Case 1, the surgery was radical and no adjuvant treatment was indicated. The patient is kept under surveillance using endoscopic examination and regular CT/ultrasonography of the chest and peritoneal cavity. At the time of writing, no evidence of the disease has been registered 65 months after the resection.

DISCUSSION

LGFMS is a soft tissue neoplasm with an unclear incidence, although, a study from western Denmark estimates the incidence is 0.18 per million\(^6\). The disease preferen-
LGFM

tially affects young and middle-aged adults with a median age ranging from 29 to 40 years. Nevertheless, LGFM can develop at any age including extremely rare cases in infants, such as the case of a 9-year-old boy with LGFM of the superior turbinate.

The typical symptom is a painless, slow growing mass. The size of the tumor varies from 1 to approximately 19 cm at the time of the diagnosis. In our cases, tumors were 4 and 5.5 cm in their greatest diameter. Despite being quite large masses concerning location, probably the slow, noninvasive growth of the tumors caused late and only less serious symptoms. The majority of the tumors are located in the soft tissues of the trunk, lower and upper extremities, but even atypical sites such as heart or thyroid gland are described. The clinical course of the disease is unpredictable and insidious. Based on the case series with long term follow up data there is a high potential to develop late local recurrences and distant metastases. In 2011, Evans et al. presented a series of 33 cases where 21 patients had a local recurrence and 15 patients had a distant metastasis. Moreover, the median time to the development of the metastasis was 5 years, ranging from 0 to 42 years. Guillou et al. published a series of 30 patients with long term follow up data, reporting recurrence and metastasis rate to be 20%. Median time to local recurrence and metastasis were 276 and 132 months, respectively. The critical aspect is that 71.4% of patients with a follow up longer than 7 years developed a metastasis. Tang et al. in 2010 reviewed data of 273 patients with hyalinizing spindle cell tumor or LGFS, 184 of whom had follow up data available. 29% of patients had local recurrence and 18% of patients were diagnosed with a distant metastasis. These data support the necessity for a long term follow up of patients diagnosed with LGFS.

Radical surgery with wide margins is the treatment of choice. In our cases, the procedures were complicated by severe bleeding from the tumors. Especially in endoscopic endonasal surgery, bleeding is the main enemy. The reports from Manes et al. and Varsak et al. do not mention excessive bleeding despite the use of endoscopic approach. However, both the cases were much younger patients with significantly smaller tumor size and less prominent spread to the upper and dorsal parts of the nasal cavity, based on the presented MRI. Given our experience, preoperative embolisation of LGFM in this location should be considered.

On histological examination, LGFM shows bland looking spindle cell proliferation in a swirling, whorled growth pattern. Alternating myxoid and fibrous zones are characteristic. Immunohistochemically there is no specific marker for LGFM, yet the tumor shows typically strong and diffuse cytoplasmic positivity to MUC4 (ref.12). It is a transmembrane glycoprotein expressed on the epithelial surfaces of many organ systems and on some tumors such as pancreatic, biliary duct or lung adenocarcinomas. Its function is associated with phosphorylation of Erb2b tyrosinekinase that plays an important role in the proliferation and differentiation of cells. A characteristic genetic feature is balanced translocation t (7;16) (q34; p11), that leads to fusion of the FUS and CREB3L2 genes. In a minority of cases there can be a variant fusion FUSCREB3L1 or EWSR1-CREB3L1 (ref.13).

Based on a clinical examination, imaging methods and immunohistochemistry, the differential diagnosis is quite large. Pathologic conditions that can show similar patterns are myxofibrosarcoma, dermatofibrosarcoma, fibromatosis, myxoid neurofibroma or sclerosing epitheloid fibrosarcoma (SEF). Moreover, SEF is positive for MUC4 up to 78% (ref.14). There is also a tumor variant called hybrid SEF-LGFMS, that contains areas indistinguishable from LGFM and shows 100% positivity for MUC4 staining.

CONCLUSIONS

LGFS is a rare disease that even less frequently appears in the head and neck. Radical surgery is the primary treatment option although in head and neck there is a risk of serious morbidity and esthetic defect. In view of this fact, the endoscopic approach should be applied when possible. Nevertheless, endoscopic surgery can be a challenging procedure due to the possible extensive bleeding from the tumor as happened in our cases. Based on these experiences, preoperative embolisation of the tumor should be considered. However, even after successful radical surgery, prolonged follow up is necessary due to the metastasizing capacity of the tumor and the possibility of late recurrences.

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Fig. 3. Microscopic view of the tumor resected from Patient 2, 100x magnification.
evaluation and provided microscopic images of the tumor; JP: managed patient treatment and supervised manuscript preparation.

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REFERENCES