Primary oral soft tissue angiosarcoma of the cheek: a case report and literature review

Benjamin Fomete¹, Modupe Samaila², Sunday Edaibgini³, Rowlan Agbara¹, Uche Albert Okeke¹

¹Department of Maxillofacial Surgery, ²Department of Pathology, ³Division of Cardiothoracic Surgery, Department of Surgery, Ahmadu Bello University Teaching Hospital, Zaria, Nigeria

Abstract (J Korean Assoc Oral Maxillofac Surg 2015;41:273-277)

Angiosarcoma is a rare and aggressive malignant tumor that has a poor prognosis. It represents less than 1% of all malignancies occurring in the oral cavity and salivary glands. We present a 35-year-old male with angiosarcoma of the cheek following traumatic injury and a review of the current literature.

Key words: Primary, Angiosarcoma, Cheek, Oral, Soft tissue

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I. Introduction

Angiosarcoma (AS), also known as malignant hemangioendothelioma, is a rare and aggressive malignant tumor of endothelial cells of vascular or lymphatic origin. These tumors typically have a poor prognosis, and they represent less than 1% of all malignant tumors¹ and between 1% to 2% of soft tissue sarcomas²,³. About 50% to 52% of all ASs occur in the head and neck³,⁴ and are found predominantly on the scalp and face of elderly men²,⁵. One study showed that ASs represented 11.2% of all sarcomas in the the head and neck region⁶.

AS is extremely rare in the oral cavity and salivary glands, accounting for 1% of all ASs reported in the literature¹,⁵. Although the tumor can occur in any location, the most common sites are soft tissue, skin⁶, trunk, and limbs⁷.

The tumor type is represented equally between sexes³,⁷; however, Fanburg-Smith et al.⁸ suggest a mild male predominance (ratio 1.07:1).

There is inconsistency in the data on age-related tumor involvement because some authors postulate that older patients have a greater propensity for AS, while others suggest that AS favors those of a younger age⁷. In general, all age groups can be affected, though it is prevalent among older white men and often presents as a cutaneous lesions¹. The median age of those who develop AS, according to Yang et al.¹, is 60 to 70 years, and the mean age, according to Fanburg-Smith et al.⁸, is 55 years. Sun exposure has been implicated in the etiology of this tumor². ASs are highly aggressive and can arise in association with some recognized clinical conditions, such as chronic lymphedema and previous radiotherapy, or they can arise de novo². AS is initially inconspicuous and is often diagnosed as a simple bruise. It expands with gradual centrifugal infiltration, eventually resulting in nodule formation and skin ulceration². AS presents as an extremely aggressive lesion with propensity for local recurrence and distant metastasis, leading to poor prognosis³.

The overall prognosis of AS is poor, even compared with that of other head and neck sarcomas². The five-year survival in the head and neck is 10% to 35%³,⁴, and approximately half of AS patients die within 15 months of diagnosis⁵. Early detection and treatment are essential to control this highly malignant soft tissue cancer⁶.

We present a case of a 35-year-old male with primary AS of the cheek initially misdiagnosed as sclerosing hemangioma and a review of relevant literature.
II. Case Report

A 35-year-old male was seen at the maxillofacial surgery clinic of our hospital with a four-year history of left facial swelling of insidious growth. He reported a history of assault by armed robbers who hit him with a gun on the left cheek, where he developed swelling eight months later. Three months prior to his presentation at our center, he presented at a peripheral hospital where he underwent a surgical procedure and received a histopathological report of sclerosing hemangioma.

At presentation, the left cheek mass was covered with a dressing (Fig. 1), which on exposure revealed an ulcerated mass with no discharge or bleeding. The mass measured 10 cm by 7 cm and was firm to the touch. It was attached to underlying structures and protruded intraorally with indentation of the overlying mucosa. A computerized tomographic angiogram of the cheek (Fig. 2) revealed a rounded, isodense and brightly enhancing mass over the body and anterior to the ramus of the left hemimandible. The enhancement was the same density as the adjacent left facial artery, which coursed through the medial aspect of the mass. A diagnosis of a left facial artery pseudo-aneurysm was made. Other tests such as full blood chemistry, electrolyte count, creatinine, and random blood glucose were within the normal range for his age and sex.

He was prepared for surgery, and on the operating table, the feeder vessel was ligated (Fig. 3) to prevent torrential bleeding from the presumed pseudoaneurysm or sclerosing hemangioma. The mass was surgically removed via an extraoral approach under general anesthesia. The whole specimen (Fig. 4) was sent to the histopathology laboratory of our hospital and was reported as AS. (Fig. 5)

He had an uneventful recovery and was transferred to the oncology team for further management on the seventh postoperative day. His management was interrupted by a health worker strike, and he was discharged home. He was seen at the clinic for follow-up three months later. No obvious or residual tumour was seen, and he declined further medical intervention, including chemotherapy and radiotherapy.

The tumor specimen was in the form of two irregular shaped tissues, one of which was covered with skin and the other ulcerated, measuring 8×7.5×3 cm and 9×9×6 cm, re-
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The etiology of AS is not fully understood, although the sun has been implicated as a causative agent in Caucasian patients. Chronic lymphedema and irradiation have been described as the most important etiological factors in the pathogenesis of AS. Only about 10% of ASs are associated with chronic lymphedema, even though it is the most widely accepted sign of AS in skin and soft tissue lesions. Radiation-induced AS usually occurs in the irradiated fields many years after radiotherapy, with a median time of 8.6 years having been reported. AS is also said to occur in pre-existing or intermediate vascular lesions. Several cases of AS arising spontaneously from hemangiomas or vascular malformation have been reported, but the mechanism is not clear. Even though most authors think trauma can only alter an already existing lesion, it has been reported as an

![Fig. 3. Isolation of major vessels.](image1)

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![Fig. 4. Excised specimen cut open.](image2)

**Fig. 4. Excised specimen cut open.**
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![Fig. 5. A. Intraluminal buds of tumor cells (H&E staining, ×40). B. Sheets of epithelioid tumour cells (H&E staining, ×40). C. Strong positivity with highlights of neoplastic vascular channels (CD31 staining, ×40).](image3)

**Fig. 5. A. Intraluminal buds of tumor cells (H&E staining, ×40). B. Sheets of epithelioid tumour cells (H&E staining, ×40). C. Strong positivity with highlights of neoplastic vascular channels (CD31 staining, ×40).**
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respectively, together weighing 179 g. The cut section revealed grey, white, solid, and hemorrhagic nodular surfaces.

Histology sections showed keratinized pigmented skin overlying a fibrocollagenized dermis and a malignant spindle cell tumor growing in sheets, with anastomosing dilated vascular channels. The cells comprising the tumor were plump and round to spindle-shaped with hyperchromatic to vesicular nuclei, prominent nucleoli, and scanty cytoplasm. There was minimal fibrocollagenized stroma. Other areas showed hemorrhage, fibrin deposits, and necrosis.

**III. Discussion**

AS is said to account for about 2% of all soft tissue sarcomas, while oral AS is very rare. It has a predisposition for the skin and superficial soft tissue. The etiology of AS is not fully understood, although the sun has been implicated as a causative agent in Caucasian patients. Chronic lymphedema and irradiation have been described as the most important etiological factors in the pathogenesis of AS. Only about 10% of ASs are associated with chronic lymphedema, even though it is the most widely accepted sign of AS in skin and soft tissue lesions. Radiation-induced AS usually occurs in the irradiated fields many years after radiotherapy, with a median time of 8.6 years having been reported. AS is also said to occur in pre-existing or intermediate vascular lesions. Several cases of AS arising spontaneously from hemangiomas or vascular malformation have been reported, but the mechanism is not clear. Even though most authors think trauma can only alter an already existing lesion, it has been reported as an
etiological factor. Fanburg-Smith et al. reported 29 cases with 22 primary AS lesions in the tongue (n=9), parotid gland (n=4), lip (n=4), submandibular gland (n=3), soft palate (n=1), and hard palate (n=1) and seven secondary lesions in the gingivae (n=4) and parotid gland (n=3). The tongue, parotid gland, and lip are commonly involved in adults.

Our case occurred in the cheek. AS can arise from either a vascular or lymphatic origin, and predisposing factors to its development include irradiation, trauma, and advanced age. Nagata et al. reported that 35 cases have been documented in the English literature.

Our patient reported a definite history of trauma, although the etiology and triggering factors are largely unknown in many cases.

AS has been categorized into five broad groups based on a wide range of associated clinical scenarios: lymphedema-associated AS, radiation-induced AS, post-breast cancer AS, soft tissue AS, and cutaneous AS. AS of the oral cavity can involve oral soft tissue, minor salivary glands, or bones. Our patient’s AS affected the oral soft tissue of the cheek. Several reports of primary AS involving the oral soft tissue, lip, palate, tongue, parotid gland, and submandibular glands are also documented. The clinical presentation is often a mass and/or bleeding, pain, bruising, persistent hematoma, edema and ulceration, or round or ovoid nodules that appear red to bluish in color.

The clinical differential diagnoses include pyogenic granuloma, hemangioma, hemangioendothelioma, Kaposi’s sarcoma, and ulcerative squamous cell carcinoma. Our working diagnosis in this case was sclerosing hemangioma based on the histology report and the computed tomography (CT) scan the patient brought from another center. Primary ASs are often misdiagnosed as other lesions because of their rarity. Our case was initially reported as sclerosing hemangioma, and while a CT scan supported a facial artery aneurysm, other missed diagnoses include pyogenic granuloma. Delayed or missed diagnosis is associated with poor treatment outcome.

According to Nagata et al., AS can show a wide spectrum of histological differentiation, from a well differentiated neoplasm showing anastomosing vascular channels lined by atypical endothelial cells with low mitotic activity to poorly differentiated tumors without prominent vasoformative activity, composed of solid sheets of epithelial-like or spindle cells. Fanburg-Smith et al. stated that the most common morphologic type of AS is solid spindled vasoformative, but that almost one-third of oral and salivary gland ASs are the rare epithelioid variant.

Immunohistochemical analysis is essential for differentiating AS from similar lesions such as hemangioendothelioma, hemangiopericytoma, epithelioid sarcoma, Kaposi’s sarcoma, perivascular epithelioid tumor (malignant Pecoma), spindle cell carcinoma, malignant melanoma, intravascular endothelial hyperplasia, epithelioid AS, and lymphangiosarcoma.

Positive staining for factor VII and Ulex europaeus agglutinin has been used to diagnose vascular pathology. CD31, a membrane glycoprotein, is the best vascular marker of endothelial phenotypes, particularly in poorly differentiated AS, where factor VIII and U. europaeus are negative. Positive staining for thrombomodulin and E-selectin also support the diagnosis of AS. Tumors showing vasoformative channels and intracytoplasmic vacuoles on hematoxylin and eosin sections are likely angiogenic. Positive reactions to F-VIII-RA, CD34, and CD31 endothelial markers indicate endothelial characteristics. The presence of red blood cells in the vasoformative channels suggests that the tumor is not a lymphatic tumor but a vascular tumor.

Surgery is the main stay of treatment, and local primary tumors can be effectively excised. However, there is a high associated rate of local recurrence and distant metastasis, which negatively impacts patient survival. According to Köhler et al., the lesion must be excised with adequate three-dimensional surgical margins, which can be difficult to achieve in the head and neck region.

A combined modality of treatment, including postoperative radiotherapy, is recommended. The general outcome of primary AS of the oral cavity is good, with a mean survival period of 7.3 to 7.6 years, compared to primary cutaneous and deep soft tissue tumors. Our patient underwent surgery and declined oncological treatment. Because most patients present with systemic metastasis, chemotherapy is often involved in the treatment of AS. Complete remission of AS has been reported with the use of liposomal doxorubicin alone or in association with radiotherapy. Though there was no evidence of residual disease on follow-up, the risk of recurrence without further adjunct therapy cannot be overlooked. Due to its rarity, AS is associated with no significant or clearly definitive prognostic factors. Lesion size has been used as the only significant predictor, while histology has had no impact. Additionally, the lesion site was considered a significant prognostic factor as tumors in the scalp had a worse prognosis. However, Köhler et al. found...
that grade was the most important prognostic factor.

Metastatic spread is usually hematogenous, and a common site of involvement is the lungs. Lymphatic spread is reported in less than 20% of ASs involving the head and neck region. In our case, no lymph node involvement or distant metastases were observed at initial presentation or during the follow-up period; however, we did not carry out any lymph node survey.

Primary AS of the cheek is rare, and its diagnosis and treatment still pose a challenge in clinical settings.

Conflict of Interest

No potential conflict of interest relevant to this article was reported.

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