CHARACTERIZATION OF ALVEOLAR SOFT PART SARCOMA OF THE TONGUE: A CLINICO-PATHOLOGIC STUDY AND SCOPING REVIEW

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ABSTRACT

Background: Alveolar soft part sarcoma (ASPS) is a rare malignant soft tissue tumour. There is a dearth of literature analyzing its features on the tongue.

Objectives: This study aims to conduct a scoping review to describe the essential clinico-pathologic features, treatment modalities and outcome of previously reported tongue ASPS (TASPS) and new cases at our center.

Methods: A search of databases (PubMed, Medline, Cochrane and Google Scholar) and the internet for articles on TASPS written in English was conducted. Information extracted included clinico-pathological and demographic data. Descriptive statistics was used for analysis.

Results: A total of 49 articles were eligible for this study. In all, 81 cases were utilized. Asian studies accounted for most cases 35(43.2%) and a slight female preponderance of 1.1 was seen. Most cases - 38 (46.9%), occurred in the 1st decade and the base of tongue was the most common location in 19 (39.6%) cases. Also, tumour metastasis was present in 14 (25.9%) cases. Transcription Factor E3 (TFE3) – 8 (24.2%) and Neuron Specific Enolase (NSE) – 8 (24.2%) were the most common immunohistochemical stains used and were both expressed 7 out of 8 cases (87.5%). Most common treatment modality was surgery and 42 (82.4%) cases managed by surgery alone were free of disease at ≤ 5 years of follow up.

Conclusions: TASPS slightly affected the female gender and tongue base more commonly. It occurred more in the first two decades of life. Use of standard investigative tools for management will allow for better appraisal of research findings.

Keywords: Tongue; Alveolar; Soft-part; Sarcoma; Treatment outcome

INTRODUCTION

Alveolar soft part sarcoma (ASPS) is a rare malignant soft tissue tumour that accounts for about 1% of all soft tissue sarcomas.\(^1\) It was first described in 1952 by Christopherson and Stewart.\(^3\) Despite numerous studies since then, the histogenesis, biologic behavior and best treatment modality has remained debatable. The head and neck region is the favored site for ASPS in children and adolescents, while the thigh and buttocks are common sites for ASPS in adults.\(^4\) Also, a female predilection has been reported in cases occurring in the 1st and 2nd decades of life while a slight male preference was observed after the 3rd decade.\(^7\)

Presentation of ASPS is usually that of a slow growing painless mass, with a high rate of metastasis to the lungs, bone, and the brain, which could occur long after excision of the primary tumour.\(^7\) ASPS could present clinically as a vascular lesion and magnetic resonance imaging (MRI) of the tumour with contrast enhancement is ideal to demonstrate its features on the tongue. Microscopically, ASPS consists of large polygonal to round cells with distinctive cell membrane, abundant eosinophilic granular cytoplasm, round to oval eccentric nuclei with prominent nucleoli which may be multiple. Neoplastic cells are characteristically disposed in nested or organoid growth pattern separated by thin fibrous septa.\(^6\) The cells may appear non-cohesive, giving it the alveolar pattern. Those without organoid patterns have also been described as well as those with clear cytoplasm. The solid pattern is more frequently seen in pediatric cases.\(^15\) The tumour is well vascularized by delicate sinusoidal vascular channels lined by a single layer of endothelial
cells. Pleomorphism and mitosis are infrequent. About 80% of ASPS have intracytoplasmic, periodic acid–schiff positive, diastase-resistant rhomboid- or rod-shaped crystals.16

Furthermore, ASPS have been reported to commonly occur on the tongue in many studies, as well as in case reports and constitute 25% of all ASPS.17-21 Also, tongue alveolar soft part sarcoma (TASPS) occurs in patients much younger than those for ASPS from other anatomical locations particularly in females.2,18,22,23

There are many reports describing the clinico-pathologic features of ASPS.10,18,24 However, there is a dearth of literature analyzing these features in tongue tumours only, despite the tongue being a common site of presentation in the head and neck region. Therefore, it is desirable to assess the characteristics of TASPS and to assess the available treatment modalities necessary to achieve a desirable outcome in the management of this entity. This study aims to describe the essential clinico-pathologic features, treatment modalities and outcome of previously reported TASPS by conducting a scoping review along with present cases seen at the Oral Pathology Department, University College Hospital, Ibadan.

MATERIALS AND METHODS

Study design

This study was a review of previous studies describing the clinico-pathologic features of TASPS. A scoping review was conducted because the available studies on TASPS varied in their methods and data, thus precluding the conduct of a meaningful meta-analysis. The review was performed following the Preferred Reporting Items for Systematic Reviews and Meta-Analyses extension for Scoping Reviews (PRISMA-ScR) Checklist.25

Methods

The histopathology records of the Department of Oral Pathology, University College Hospital (UCH), Ibadan over a period spanning 26 years were examined. All entries of cases diagnosed as ASPS were retrieved, while TASPS were identified for analysis. The haematoxylin and eosin (H&E) slides of these cases were retrieved and reassessed to verify the diagnosis. Case files of eligible cases were retrieved and information on bio data, duration of symptoms, symptoms on presentation, presence of tumour spread or metastasis at diagnosis, site of metastasis if present, clinical impression, treatment received, duration of follow up and status at follow up were obtained, cases with incomplete records were excluded from the study.

Literature search

To identify relevant studies, an all-inclusive search of the databases (PubMed, Medline, Cochrane and Google Scholar) as well as an internet search of articles written in English language was conducted between August and October 2019. Keywords used for the search included a blend of “alveolar soft-part sarcomas (ASPS),” “soft-tissue tumours,” and/or “tongue.” Also, relevant citations identified in the reference lists of selected articles were included in the search. The search lists from the electronic sources were merged and duplicates were removed. The title and abstract of the identified articles were screened to remove studies outside the scope of this review after which the full text of all potentially eligible articles were retrieved for further analysis. Articles that did not satisfy the inclusion criteria were excluded from further consideration. Also, a manual search of bibliographies of identified articles was done by cross referencing eligible publications on ASPS from 1957 till date while relevant citations identified in the reference lists of selected articles were included in the search to identify additional studies of interest. The selection process is displayed in a flow chart (Figure 1).

Criteria for eligibility

Articles included were human case reports/case series, letter to editors and review articles on ASPS of the tongue either in whole, or as part of a series on ASPS. Articles that were not available in English or which the full text could not be obtained were excluded from the study. Similarly, ASPS that metastasized to the tongue as well as cases with incomplete data were excluded from the study.

Extraction of data

A proforma was used to extract data from eligible articles by two investigators (AOA, BFA) independently. Information extracted included clinico-pathological and demographic data such as year of publication, country of publication, type of study, patients’ age, gender, location and surface of tongue affected, duration of symptoms, symptoms on presentation, presence of tumour metastasis at diagnosis, site of metastasis if present, clinical impression, result of immunohistochemical studies (when available), treatment type, duration of follow up and status of patients at last follow up. Any inconsistencies were resolved by consensus with a third investigator (GOO).

Statistical analysis

Descriptive statistics was used for analysis. Relevant data were extracted from the included studies and variables were presented using summary statistics and tables. Data analysis was done using SPSS software version 21 (IBM Corporation, Armonk, NY, USA).
RESULTS
Initial electronic search of the databases identified 29 potentially eligible articles. An additional 31 publications were identified from other sources (bibliography of initially identified articles). After initial review of the titles and abstracts, four duplicate articles were removed and 56 articles which met the inclusion criteria were identified. Eight of these were excluded because two were not written in English language, while full text articles were unobtainable for another six. Also, one eligible article was identified and included following a hand search. Thus, a total of 49 articles (39 case reports and 10 case series) were utilized in this study.

Furthermore, five cases of ASPS were identified from the records of Oral Pathology Department, UCH, Ibadan. Four affected the tongue, while one affected the cheek and was excluded from further analysis. The age range of cases was 6 to 34 years; while male to female ratio was 3:1. Also, the site of predilection was the dorsal surface of the anterior tongue (Figure 2). Duration of symptoms ranged from three months to four years with tumours in the anterior dorsum and sulcus terminalis areas having a shorter duration than posterior and ventral tongue tumours.

Histopathology of all cases showed tissue disposed in organoid pattern of large round to oval eosinophilic cells separated by moderately to highly vascularized fibrous connective tissue stroma. Individual cells have abundant granular cytoplasm with some having eccentric nuclei while others had vesicular nuclei, (Figure 3). The clinical data of the present cases have been summarized in Table 1.

Table 1: Characterization of TASPS in Ibadan.

| Case | Gender | Age | Site               | Surface | Duration | Size (cm) | Treatment | Follow up | Status at follow up |
|------|--------|-----|--------------------|---------|----------|-----------|-----------|-----------|-------------------|
| 1    | Male   | 29  | Sulcus terminalis | Dorsum  | 3 months | NR        | NR        | NR        | LTFU              |
| 2    | Male   | 6   | Anterior           | Dorsum  | 3 months | 6         | Surgery   | 13 months | FOD               |
| 3    | Female | 34  | Anterior           | Dorsum  | 6 months | 5         | None      | 19 months | DOD               |
| 4    | Male   | 17  | Posterior          | Dorsum  | 48 months| NS        | None      | 12 months | AWD               |

NR – No record; LTFU – Lost to follow up; FOD – Free of disease; DOD – Died of disease; AWD – Alive with disease

In all, 77 cases from 49 articles and four cases from records of Oral Pathology Department, UCH, Ibadan (totaling 81 cases) were used. Table 2 shows a list of the publications and the number of cases. Asian studies accounted for 35 (43.2%) cases, while North American and European studies recorded 25 (30.9%) and 12 (14.8%) cases respectively (Figure 4). There was a slight female preponderance of 1.1 and 33 (36.4%) followed by granular cell myoblastoma/tumour 6/33 (18.2%) and dermoid cyst 2/33 (6%). Only 54/81 (66.7%) cases recorded the presence or absence of tumour metastasis either at presentation or at any point during treatment. Tumour metastasis was present in 14/54 (25.9%) cases, while it was not seen in 40/54 (74.1%) cases. Also, the most common site of tumour metastasis were regional lymph nodes
### Table 2: Characteristics of reviewed literature of TASPS.

| S/N | Year | Author(s)          | Country | Age & Gender | Site & Surface | Size (cm) | Treatment | Follow up (Months) | Status at follow-up |
|-----|------|--------------------|---------|--------------|---------------|-----------|-----------|-------------------|---------------------|
| 1   | 1952 | Christopherson et al. | USA     | 12/F         | Dorsun        | 5 cm (Gross) | Surgery   | 60 months         | NED                 |
| 2   | 1979 | Spector et al.      | USA     | 17/F         | Base          | 48 cm      | Surgery, RTH, Chemotherapy, | 60 months | DOD                 |
| 3   | 1983 | King and Fee        | USA     | 5/F          | Anterior      | 1 cm       | Surgery   | 24 months         | FOD                 |
| 4   | 1984 | Komori et al.       | Japan   | 11/F         | Base          | 26 cm      | Surgery   | 60 months         | FOD                 |
| 5   | 1984 | Chaudhry et al.     | Pakistan| 0.3/F        | Dorsun        | 2 cm       | Surgery   | 60 months         | FOD                 |
| 6   | 1985 | Sawyer et al.       | Nigeria | 16/F         | Base/Ventral  | 1 cm       | Surgery   | 24 months         | FOD                 |
| 7   | 1987 | Donald              | USA     | 15/F         | Torgue NOS    | -          | Surgery, RTH | 42 months | FOD                 |
| 8   | 1989 | Simmons et al.      | USA     | 1.4/F        | Dorsun        | 15 cm      | Surgery, RTH | 12 months | FOD                 |
| 9   | 1989 | Çetik et al.        | Turkey  | 13/F         | Dorsun        | 15 cm      | Surgery, RTH | 12 months | FOD                 |
| 10  | 1990 | Takita et al.       | Japan   | 15/M         | Dorsun        | 28 cm (Scar) | Surgery, Chemotherapy | 36 months | FOD                 |
| 11  | 1990 | Matsuno et al.      | Japan   | 6/M          | Torgue NOS    | 3 cm       | Surgery, Chemotherapy | 27 months | AWD                 |
| 12  | 1993 | Carson et al.       | USA     | 64/M         | Base          | 8 cm       | Chemotherapy, RTH | 36 months | DOD                 |
| 13  | 1993 | Ooi et al.          | Singapore | 21/M     | Anterior-base/ Dorsun | 4 cm | Surgery | 4 months | FOD                 |
| 14  | 1998 | Hunter et al.       | USA     | 3/F          | Torgue NOS    | -          | Surgery   | 48 months         | FOD                 |
| 15  | 1999 | Bentley et al.      | UK      | 5/F          | Base          | 65 cm      | Surgery, RTH | 42 months | FOD                 |
| 16  | 2000 | Casanova et al.     | Italy   | 18/F         | Torgue NOS    | -          | Surgery   | 12 months         | FOD                 |
| 17  | 2000 | Casanova et al.     | Italy   | 5/F          | Torgue NOS    | -          | Surgery, Chemotherapy, Surgery | 129 months | FOD                 |
| 18  | 2000 | Yoshida et al.      | Japan   | 2/F          | Dorsun        | 2 cm       | Surgery   | 86 months         | FOD                 |
| 19  | 2003 | Aiken and Stone     | USA     | F/34         | Base          | -          | Surgery, Chemotherapy | 24 months | MD                  |
| 20  | 2004 | do Nascimento Souza et al. | Brazil | 1.5/F        | Lateral/Dorsun | 3 cm | Surgery, Chemotherapy | 60 months | FOD                 |
| 21  | 2004 | Fanburg-Smith et al. | USA     | 3/F          | Lateral       | 25 cm      | Surgery, Chemotherapy | 122 months | NED                 |
| 22  | 2004 | Fanburg-Smith et al. (Castle 1999) | USA     | 3/M          | Lateral       | 08 cm      | Surgery | 48 months         | NED                 |
| 23  | 2004 | Fanburg-Smith et al. (2) | USA     | 3/F          | Lateral       | -          | Surgery   | 120 months        | NED                 |
| 24  | 2004 | Fanburg-Smith et al. (4) | USA     | 5/F          | Base          | -          | Surgery   | 122 months        | FOD                 |
| 25  | 2004 | Fanburg-Smith et al. (5) | USA     | 5/F          | Lateral       | 5 cm       | Surgery   | 37 months         | NED                 |
| 26  | 2004 | Fanburg-Smith et al. (6) | USA     | 5/F          | Mid portion   | 1 cm       | Surgery   | 264 months        | FOD                 |
Table 2: Cont'd

| No. | Year | Authors et al. | Country | Gender | Laterality | Diameter | Treatment | Follow-up | Site | Tumor spread |
|-----|------|----------------|---------|--------|------------|----------|-----------|-----------|------|--------------|
| 27  | 2004 | Fanburg-Smith et al. (7) | USA | 5/F | Lateral | - | Surgery | - | - | - |
| 28  | 2004 | Fanburg-Smith et al. (8) | USA | 6/N | Tongue NCS | - | Surgery | - | 336 months | FOD |
| 29  | 2004 | Fanburg-Smith et al. (9) | USA | 6/N | Mid-posterior | 2.5cm | Surgery | - | 324 months | FOD |
| 30  | 2004 | Fanburg-Smith et al. (10) | USA | 7/N | Ventral | 1.3cm | Surgery | - | - | - |
| 31  | 2004 | Fanburg-Smith et al. (11) | USA | 7/N | Lateral | 2.5cm | Surgery, Chemotherapy | 192 months | NED |
| 32  | 2004 | Fanburg-Smith et al. (12) | USA | 11/M | Base | 2.5cm | Surgery | - | 300 months | NED |
| 33  | 2004 | Fanburg-Smith et al. (13) | USA | 20/M | Base | 1cm | Surgery | - | - | - |
| 34  | 2004 | Fanburg-Smith et al. (14) | USA | 21/F | Lateral | 3cm | Surgery | - | - | - |
| 35  | 2005 | Kim et al. (1) | Korea | 10/M | Tongue NCS | - | Surgery | - | 6 months | FOD |
| 36  | 2005 | Kim et al. (2) | Korea | 4/F | Tongue NCS | - | Surgery | - | 8 months | FOD |
| 37  | 2005 | Kanhere et al. (1) | India | 6/N | Tongue NCS | 1.5cm | Surgery | - | 18 months | FOD |
| 38  | 2005 | Kanhere et al. (2) | India | 42/M | Tongue NCS | 3.5cm | Surgery, RTHI | 21 months | RD |
| 39  | 2005 | Kanhere et al. (3) | India | 12/M | Tongue NCS | 4.5cm | Surgery | - | 24 months | FOD |
| 40  | 2005 | Kanhere et al. (4) | India | 43/M | Tongue NCS | 2.5cm | Surgery, RTHI | 72 months | AWD |
| 41  | 2006 | Carvao-Silva et al. | Brasil | 17/F | Anterior/Dorsum | 2cm | Surgery | - | 12 months | FOD |
| 42  | 2006 | Rua et al. | Korea | 3/F | Lateral/Dorsum | 2cm | Surgery | - | 32 months | FOD |
| 43  | 2007 | Raghunandhan et al. | India | 13/F | Base | 2.5cm | Surgery, RTHI | 36 months | FOD |
| 44  | 2008 | Tapsiz et al. | Turkey | 18/F | Dorsum | 4.5cm | Chemotherapy | - | - | - |
| 45  | 2009 | Rodriguez-Velasco et al. | Mexico | 2/F | Lateral | 1.5cm | Chemotherapy | - | 34 months | FOD |
| 46  | 2009 | Baghram et al. | Turkey | 18/F | Base | 6cm | Surgery, RTHI | 10 months | DOD |
| 47  | 2010 | Nousios et al. | Greece | 2/M | Dorsum | 3.5cm | Surgery | - | 42 months | FOD |
| 48  | 2010 | Kumar et al. | India | 7/N | Lateral/Dorsum | 3cm | Surgery, Chemotherapy | - | 11 months | FOD |
| 49  | 2010 | Floy et al. | UK | 24/M | Lateral | 1cm | Surgery | - | 12 months | FOD |
| 50  | 2011 | Asharaz et al. | India | 75/M | Anterior/Dorsum | 3cm | Surgery | - | 36 months | FOD |
| 51  | 2011 | Conice et al. | Spain | 5/F | Base/Ventral | 4cm | Surgery, Chemotherapy | - | 36 months | FOD |
| 52  | 2012 | Rekhi et al. (1) | India | 24/F | Tongue NCS | - | Surgery | - | - | - |
| 53  | 2012 | Rekhi et al. (2) | India | 18/F | Tongue NCS | - | Surgery | - | - | - |
| 54  | 2013 | Agyris et al. | Greece | 4/N | Sulcus terminalis/ Dorsum | 2cm | Surgery | - | 7 months | FOD |
| 55  | 2013 | Adeyemi et al. | Nigeria | 27/M | Ventral | 6cm | Surgery | - | - | - |
| 56  | 2014 | Kingo et al. | India | 14/M | Anterior/Dorsum | 4cm | Surgery | - | - | - |
| 57  | 2014 | Meng et al. | China | 4/N | Root | 4.5cm (CT) | Chemotherapy, RTHI | 30 months | FOD |
Table 2: Cont’d

| Year | Authors | Country | Stage | Tumor Location | Tumor Size | Treatment | Follow-up | Outcome |
|------|---------|---------|-------|----------------|------------|-----------|-----------|---------|
| 54   | Liu et al. | Taiwan | 27/F | Ventral and FOM | 5 cm | Surgery, RTH | 24 months | FOD |
| 55   | Wang et al. | China | 20/F | Base | 2.5 cm | Surgery | 34 months | NED |
| 56   | Wang et al. | China | 19/F | Base | 3.5 cm | Surgery | 10 months | NED |
| 57   | Wang et al. | China | 11/M | Dorsal | 3 cm | Surgery | 14 months | NED |
| 58   | Wang et al. | China | 14/F | Ventral and FOM | 5 cm | Surgery | 15 months | NED |
| 59   | Wang et al. | China | 28/F | Ventral and FOM | 6 cm | Surgery, RTH | 21 months | AWD |
| 60   | Wang et al. | China | 4/M | Ventral and FOM | 6.5 cm | Surgery | 23 months | NED |
| 61   | Wang et al. | China | 7/M | Base | 4 cm | Surgery | 29 months | NED |
| 62   | Wang et al. | China | 16/F | Base | 3.3 cm | Surgery | 35 months | NED |
| 63   | Wang et al. | China | 10/M | Base | 3.5 cm | Surgery | 77 months | NED |
| 64   | Wang et al. | China | 6/F | Ventral and FOM | 5 cm | Surgery | 60 months | NED |
| 65   | Chopra and Tanveer | India | 35/M | Lateral | 2 cm | Chemotherapy, Surgery | - | - |
| 66   | Yoshihira et al. | Japan | 23/M | Tongue NOS | - | Surgery, Chemotherapy | 4 months | MD |
| 67   | Chauve et al. | India | 8/F | Base | 3 cm | Surgery | - | LTFU |
| 68   | Kata et al. | USA | 3.8/M | Ventral | 2 cm | Surgery | 60 months | FOD |
| 69   | Eulle et al. | UK | 0.9/F | Anterior | 1.6 cm (MRI) | Surgery | 14 months | FOD |
| 70   | Leszcynska et al. | USA | 6/M | Dorsum | 1.2 cm (CT) | Surgery | 36 months | FOD |
| 71   | Hsu et al. | Taiwan | 3/M | Posterior/Dorsum | 1.2 cm | Surgery | 1 month | FOD |
| 72   | Alegria-Landa et al. | Spain | 53/M | Anterior | 2 cm | Surgery | - | - |
| 73   | Fouad et al. | Morocco | 13/M | Tongue NOS | 7.4 cm (Gross) | Surgery, Chemotherapy | 12 months | DOD |
| 74   | Present case 1 | Nigeria | 29/M | Sulcus terminalis/Dorsum | - | Surgery | - | LTFU |
| 75   | Present case 2 | Nigeria | 6/M | Anterior/Dorsum | 6 cm | Surgery | 13 months | FOD |
| 76   | Present case 3 | Nigeria | 34/F | Anterior/Dorsum | 5 cm | No treatment | 12 months | DOD |
| 77   | Present case 4 | Nigeria | 17/M | Posterior/Dorsum | - | No treatment | 16 months | AWD |

**NB:** No evidence of disease (NED) was analyzed as free of disease (FOD) for standardization. Cases reported by Fanburg-Smith et al., that were recorded as “Alive” were assumed to be free of disease except they were specified as “Alive with disease”.

M, Male; F, female; NOS, not otherwise specified; RTH, radiotherapy; FOM, floor of mouth; FOD, free of disease; NED, no evidence of disease; AWD, alive with disease; MD, metastatic disease; RD, recurrent disease; DOD, died of disease; LTFU, lost to follow-up; - no data.
Table 3: Age distribution of cases in decades.

| Years in decade | Frequency | Percentage |
|-----------------|-----------|------------|
| 0-9             | 38        | 46.9       |
| 10-19           | 24        | 29.6       |
| 20-29           | 12        | 14.8       |
| 30-39           | 3         | 3.7        |
| 40-49           | 2         | 2.5        |
| 50-59           | 1         | 1.2        |
| 60-69           | 1         | 1.2        |
| **Total**       | **81**    | **100.0**  |

Table 4: Treatment modalities of TASPS cases.

| Treatment type                                      | Frequency | Percentage |
|-----------------------------------------------------|-----------|------------|
| Surgery only                                        | 51        | 63.0       |
| Surgery and chemotherapy                            | 11        | 13.6       |
| Surgery and radiotherapy                            | 6         | 7.4        |
| Surgery, chemotherapy and radiotherapy              | 5         | 6.2        |
| No treatment                                        | 2         | 2.5        |
| Chemotherapy only                                   | 2         | 2.5        |
| Chemotherapy and radiotherapy                       | 1         | 1.2        |
| Chemotherapy, then surgery                          | 1         | 1.2        |
| Chemotherapy, radiotherapy, surgery and brachytherapy| 1         | 1.2        |
| Not specified                                       | 1         | 1.2        |
| **Total**                                           | **81**    | **100.0**  |

and the lungs, both recording 6/14 (42.8%) and 4/14 (28.6%) respectively. The lungs and the liver as well as the lungs and lymph nodes were affected in one case each, while disseminated disease occurred in 2/14 (14.3%) cases.

Immunohistochemical studies became more established in the last two to three decades of this review and were performed in 37/81 (45.6%) cases in which 33/37 (89.2%) cases were stained using the following antibodies: Transcription Factor E3 (TFE3) - (8/24.2%); Neuron Specific Enolase (NSE) - (8/24.2%); desmin (7/21.2%); actin (5/15.2%) and vimentin (5/15.2%) while four cases stained negative to all the immunohistochemical stains used. Interestingly, TFE3 and NSE were both expressed 7
Figure 2: Clinical pictures of 6 year old male, A and B shows dorsal swelling of the tongue, C shows surgical specimen and D shows tongue one-week post operatively.

Figure 3: Histopathology of TASPS cases in Ibadan. Photomicrograph shows (A) - solid pattern having tissue disposed in organoid arrangement, separated by vascularized fibrous septae H & E X 40; (B) - shows large oval to round eosinophilic cells H & E X 100 and (C) – shows non-cohesive individual cells having abundant granular cytoplasm H & E X 400
out of 8 cases (expression rate of 87.5%) while desmin, actin and vimentin were expressed 5 out of 7 (71.4%), 4 out of 5 (80%) and 3 out of 5 (60%) cases respectively. Polymerase chain reaction was also used in three instances to detect the presence of TFE3.

Most common treatment modality was surgery in 51 (63%) cases, followed by surgery + chemotherapy in 11 (13.6%) cases and surgery + radiotherapy in 6 (7.4%) cases (Table 4). Subsequently, 42/51 (82.4%) cases that were managed by surgery alone were free of disease at ≤5 years of follow up while 9/51 (17.6%) were free of disease at >5 years of follow up. All the patients (4 cases) that had follow up for over 300 months, who had surgery alone, had either “no evidence of the disease” or “were free of the disease” at the last follow up.

**DISCUSSION**

The present study presents an effort to characterize TASPS by giving a synopsis of its clinico-pathologic features from when it was first reported till date and present cases seen at our center. Our major findings include the following: Asian studies dominated the cases seen in this study, with Wang et al. contributing 10 cases having tongue involvement out of a series of 18 patients with ASPS of the oral and maxillofacial region. Increased incidence of TASPS on the Asian continent may be due to the relatively large population of Asia. Tongue ASPS slightly affected the female gender more commonly and about 76.5% of cases were diagnosed in the first two decades of life, subsequently showing a steady decline in incidence with advancing age. The base of the tongue is the most common location involved, while the dorsum is the most frequently affected surface. Interestingly, it was initially considered to be a benign lesion in some case reports and tumour metastasis occurred in 25.9% of cases that reported presence of metastasis. Also, TFE3 and NSE immunohistochemical stains had equal expression rates while surgery was the most common treatment modality.

In the present review, some findings differed from those in a previous review of 14 lingual ASPS by Fanburg-Smith et al. which to the best of our knowledge was the largest series on TASPS in English literature. Slight female preponderance was seen in the present study which differed from a male preponderance reported by Fanburg-Smith et al. Some authors have previously referred to ASPS as a disease of childhood, while some have referred to it as a disease of childhood and adolescence. In the present study, majority of ASPS occurred in the first and second decade of life but few also occurred in other age groups up to the 7th decade. Also, Fanburg-Smith et al. recorded a median age of five years while this study recorded a median age of 11 years. Similarly, the age range in this review was 11 months to 64 years while Fanburg-Smith et al. recorded a range of 3 to 21 years. The findings in this review, however concurs with the findings in a study of ASPS of the oral cavity by Shelke et al. where an age range of 1.5 – 64 years was reported. Similarly, the finding in this study on female predilection is in agreement with the outcome in the study of Shelke et al. where a female predilection for TASPS was also reported. It is probable that our larger sample size comprising of

![Figure 4: Distribution of TASPS cases according to continent.](image-url)
subjects from wider and diverse socio-cultural-geographic background may be responsible for the noted differences and may be more representative of the characteristic of TASPS.

Additionally, the findings of the base and dorsum of the tongue, as the prevalent location and surface affected in this study, were in line with the results obtained by Shelke et al. Much is yet to be understood in the preference for the tongue by ASPS in the head and neck region and the predominance of the involvement of the base as well as the dorsal surface of the tongue. The base of tongue serves as the posterior opening of the oral cavity as well as the access to the pharynx and esophagus, and the lower aspect of the nasopharynx. It is composed of sub-mucosal lymphoid tissue (lingual tonsils) and deep tongue muscles in charge of movement. Also, this region may play a role as a sump area for carcinogens and irritants. Whether a link exists between the anatomy of the base of the tongue and the preference of TASPS for this location would be a useful focus of future research. Also, most of the ASPS in this review were relatively small in size, in agreement with the previous study by Fanburg-Smith et al. Due to the location and function of the tongue, it is likely to have an early presentation with a small tumour size either due to discomfort, abnormal sensation, or interference with function which may make the patient seek help early.

ASPS like many soft tissue tumours lack specific immunohistochemical markers, which reflected in the use of a wide range of antibodies in various reports collated in this study. ASPS has been previously reported to show infrequent immunoreactivity for desmin and MyoD1 suggesting skeletal muscle differentiation. Our findings in this study revealed that various antibodies were randomly expressed in the different studies.

Nevertheless, ASPS is now believed to be a specific chromosomal alteration, der(17)t(X;17)(p11;q25), owing to the fusion of the TFE3 transcription factor gene with the alveolar soft part sarcoma critical region 1 (ASPSCHR1). The use of real-time polymerase chain reaction and fluorescent in situ hybridization in identifying fusion transcript ASPSCR1-TFE3 and TFE3 rearrangement respectively, are regarded as efficient ways for diagnosis. Similarly, this same fusion gene has been implicated in a section of translocation associated renal cell carcinomas (RCCs). However, the translocation in ASPS is unbalanced while that of translocation associated RCCs are balanced. Also, the ASPSCR1-TFE3 fusion protein plays the role of a deviant transcription factor leading to the activation of the MET signaling pathway known to stimulate angiogenesis and cell proliferation. In addition, antibodies to TFE3 exhibit nuclear positivity in ASPS; similar to findings in some translocation-associated renal cell carcinomas (RCCs), perivascular epithelioid cell neoplasm (PEComa) and granular cell tumours.

Curiously in this study, antibodies to TFE3 and NSE were used in equal number of cases and expression rate was 87.5% for each. This finding suggests that more studies would be needed to verify if NSE has a role as a reliable marker for ASPS.

Alveolar soft part sarcoma was previously reported to have a high rate of metastasis especially to the lungs, bone and the brain. However, in this review, the reported rate of metastasis for TASPS was found to be lower than expected at 25.9% of studies that reported metastasis. Also one of the cases that presented in our center who was yet to have any form of treatment, has lived with the disease for sixty months without evidence of metastasis. Adjudging that the presence of metastasis is usually seen as an indicator for malignancies, it is unclear whether TASPS represents an entity with better prognosis than ASPS in other parts of the body.

Furthermore, surgical management was the most common treatment modality employed in many studies in this review; either alone or in combination with other treatment modalities. All cases (four in all) that were followed up for over 300 months with tumour sizes ranging between 1.3 cm to 5 cm, had surgery alone and had no evidence of the disease or metastasis as at the last follow up. As opined by Fanburg-Smith et al., early diagnosis and small tumour size may be factors that influence the relatively good outcome associated with ASPS.

Study limitations

The differences in the mode of presentation of the cases posed a challenge in data retrieval and analysis since there is no uniform benchmark for case reports and series. This led to heterogeneity of results, making it challenging to pool findings from the studies included and to draw definitive conclusions from this study. Also, the cases described here may not constitute the entirety of TASPS (perhaps due to under-reporting and inaccessible full articles). However, they probably do comprise the majority of cases worldwide.

CONCLUSION

Summarily, this study has provided an up to date brief of TASPS. Tongue ASPS slightly affected the female gender more commonly and occurred more in the first two decades of life. Also, the base of the tongue was the most common location affected while surgical
management was mostly used for treatment and cases managed by surgery alone were free of disease at ≤5 years of follow up. Use of gold standard investigative tools for diagnosis and for follow up will allow for better appraisal of research findings. Longitudinal follow up of cases will also help in better understanding of this disease entity as well as the optimum treatment modality. Thus, clinicians should be suspicious of indolent appearing tongue lesions and expedite histologic assessment even when a benign lesion is suspected. This is more so when a hemorrhagic tongue swelling is being considered.

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