Peripheral Exophytic Oral Lesions: A Clinical Decision Tree

Hamed Mortazavi,1 Yaser Safi,2 Maryam Baharvand,1 Somayeh Rahmani,1 and Soudeh Jafari1

1Department of Oral Medicine, School of Dentistry, Shahid Beheshti University of Medical Sciences, Tehran, Iran
2Department of Oral and Maxillofacial Radiology, School of Dentistry, Shahid Beheshti University of Medical Sciences, Tehran, Iran

Correspondence should be addressed to Maryam Baharvand; m.baharvand@gmail.com

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1. Introduction

Lesions in the oral cavity generally present as ulcerations, red-white lesions, pigmentation, and exophytic lesions. Clinical classification of oral lesions is of great importance in the diagnostic process [1, 2]. The term oral exophytic lesions is described as pathologic growths projecting above the normal contours of the oral mucosa [2]. There are several underlying mechanisms responsible for oral exophytic lesions such as hypertrophy, hyperplasia, neoplasia, and pooling of the fluid [1], which makes it difficult to approach such lesions clinically [3, 4]. According to a national epidemiologic study by Zain et al., exophytic lesions account for 26% of all oral lesions [3]. Therefore, attempts should be done to arrive at a timely diagnosis via more logical routes like decision trees rather than test-and-error methods [3, 4]. Exophytic lesions can be classified according to their surface texture (smooth and rough), type of base (pedunculated, sessile, nodular, and dome shape), and consistency (soft, cheesy, rubbery, firm, and bony hard) [1, 4]. This narrative review paper, however, focuses on the surface shapes of the lesions as the main clinical feature in order to build a diagnostic decision tree. In this regard, oral peripheral exophytic lesions are classified as lesions with rough surface and those with smoothly contoured shape [1, 5, 6].

2. Methodology

General search engines and specialized databases including PubMed, PubMed Central, Medline Plus, EBSCO, Science Direct, Scopus, Embase, and authenticated textbooks were used by the first author and the corresponding author to find relevant topics by means of keywords such as “oral soft tissue lesion,” “oral tumor like lesion,” “oral mucosal enlargement,” and “oral exophytic lesion.” Related English-language articles published since 1988 to 2016 in both medical and dental journals were appraised. Upon compilation of data, peripheral oral exophytic lesions were categorized into two major groups according to their surface texture: smooth (mesenchymal or nonsquamous epithelium-originated) and rough (squamous epithelium-originated). Lesions with smooth surface were also categorized into three subgroups according to their general frequency: reactive hyperplastic lesions/inflammatory hyperplasia, salivary gland lesions (nonneoplastic and neoplastic), and mesenchymal lesions (benign and malignant neoplasms). In addition, lesions with rough surface were summarized in six more common lesions. In total, 29 entities were organized in the form of a decision tree in order to help clinicians establish a logical diagnosis by a stepwise progression method.
clinical trials; prospective or retrospective cohort studies), case reports, and case series on oral disease were appraised.

Out of about 150 related articles, 72 were excluded due to lack of full texts, being written in languages other than English or containing repetitive material. Finally, three textbooks and 78 papers were selected including 13 reviews, 55 case reports or case series, and 10 original articles (Figure 1). In this article, peripheral oral exophytic lesions were categorized into two major groups according to their surface texture: smooth (mesenchymal or nonsquamous epithelium-originated) and rough (squamous epithelium-originated) (Figure 2). Lesions with smooth surface were also
categorized into three subgroups according to their general frequency: reactive hyperplastic lesions/inflammatory hyperplasia, salivary gland lesions (nonneoplastic and neoplastic), and mesenchymal lesions (benign and malignant neoplasms). In addition, lesions with rough surface were summarized in six more common lesions. In total, 29 entities were organized in the form of a decision tree (Figure 3) in order to help clinicians establish a logical diagnosis by a stepwise progression method.

3. Lesions with Smooth Surface

3.1. Reactive Hyperplastic Lesions/Inflammatory Hyperplasias. Reactive hyperplasia is the most frequent phenomenon responsible for exophytic lesions in the oral cavity (Table 1). These lesions represent a reaction to some kind of chronic trauma or low grade injuries such as fractured tooth, calculus, chewing, and iatrogenic factors including overextended flange of dentures and overhanging dental restorations [7]. Reactive lesions are usually seen on the gingivae followed by the tongue, buccal mucosa, and floor of the mouth. Clinically, they appear as pedunculated or sessile masses with smooth surface. Lesions are varied from pink to red and soft to firm in terms of color and consistency [7, 8]. However, the clinical features resemble neoplastic lesions in some instances, which cause a diagnostic dilemma. The most common entities of reactive nature are pyogenic granuloma, pregnancy epulis, irritation fibroma, peripheral ossifying fibroma, peripheral giant cell granuloma, epithelium sessile, leaf-like fibroma/fibroepithelial polyp, parulis, pulp polyp, epulis granulomatous, giant cell fibroma, and inflammatory papillary hyperplasia/palatal papillomatosis [1, 4, 7].

3.1.1. Pyogenic Granuloma. Pyogenic granuloma is a common tumor-like lesion in the oral cavity appearing as a smooth or lobulated, asymptomatic mass that is usually pedunculated or sessile. The surface is characteristically ulcerated and friable, which may be covered by a yellow fibrinous membrane (Figure 4) [4, 9]. Depending on the duration of the lesion, its color ranges from shiny red to pink to purple with soft to firm in palpation. It often bleeds easily because of its extreme vascularity. Approximately in one-third of the lesions a history of trauma can be detected [10]. Pyogenic granuloma may exhibit a rapid growth, but it usually reaches its maximum size within weeks or months. The size of the lesion varies from a few millimeters to several centimeters in diameter. In 75% of all cases, the most frequently affected site is the gingivae followed by the lips, tongue, and buccal mucosa. Maxillary gingivae are more affected than mandibular gingivae, and anterior areas more

![Figure 3: Decision tree for peripheral oral exophytic lesions.](attachment:decision_tree.png)
| Entity                      | Age/Decades | Gender | Site of involvement | Surface texture | Type of base | Consistency | Color                  | Size                  | Symptom & sign               | Treatment                                      | Recurrence |
|----------------------------|-------------|--------|---------------------|-----------------|--------------|-------------|------------------------|-----------------------|-------------------------------|------------------------------------------------|------------|
| Pyogenic granuloma         | Children/young adults | Female | Gingivae            | Smooth/lobulated | Pedunculated/sessile | Soft to firm | Shiny red/pink/purple | A few mm to several cm | Asymptomatic                | Conservative surgical excision             | +          |
| Pregnancy tumor            | Pregnancy period | Female | Gingivae            | Smooth/lobulated | Sessile       | Soft to firm | Shiny red/pink/purple | A few mm to several cm | Asymptomatic                | Conservative surgical excision             | +          |
| Irritation fibroma         | 4–6 decades | Female | Buccal mucosa       | Smooth          | Pedunculated/sessile | Firm or soft | Similar to adjacent mucosa | ≤1.5 cm | Asymptomatic                | Conservative surgical excision             | Rare       |
| Peripheral ossifying fibroma | 10–19 years old | Female | Exclusively gingivae | Smooth          | Pedunculated/sessile | Firm to hard | Red to pink | <2 cm | Tooth mobility, Tooth migration, Bone loss | Surgical excision down to periosteum | 8-20%     |
| Peripheral giant cell granuloma | 5–6 decades | Female | Exclusively edentulous ridge & gingivae | Smooth | Pedunculated/ sessile/nodular | Firm | Blush purple | <2 cm or >2 cm | Bone loss, Root resorption | Complete surgical excision | +          |
| Epulis granulomatous       | >40 years   | Female | Dental socket       | Smooth          | —             | Soft to firm | Similar to adjacent mucosa | ≤1 cm to massive lesions | Asymptomatic                | Conservative treatment/surgery               | Not uncommon |
| Leaf-like fibroma          | —           | —      | Palate              | Smooth          | —             | —            | —                      | —                     | —                            | —                                              | —          |
| Epulis fissuratum          | —           | —      | Maxillary alveolar ridge | Smooth/ulcerative | Pedunculated/sessile | Firm | Pink | Up to several cm | Nontender/easily bleeding | Surgical excision & curettage | Not uncommon |
| Pulp polyp                 | Children/young adults | —      | Cervical lesions of deciduous teeth & first permanent molars | Smooth | Pedunculated/sessile | Soft to firm | Red to pink | <1 cm to large masses | Discomfort | Root canal therapy/extraction | —        |
| Giant cell Fibroma         | 3rd decade  | Female | Gingivae            | Rough surface   | Pedunculated/sessile | Firm | Pink | <1 cm (often) | Asymptomatic | Conservative surgical excision Removal of denture at night/antifungals/surgery | Rare |
| Inflammatory papillary hyperplasia | 3-5th decades | Male | Palate              | Pebbly/cobblestone | —             | Soft to firm | Red to pink | Up to several cm | Asymptomatic/symptomatic | —                                              | —          |
frequently involved than posterior areas [9–11]. There is a
female predilection and a tendency to affect children and
young adults. Patients with pyogenic granuloma are treated
by conservative surgical excision. However, recurrence is not
uncommon [9].

3.1.2. Pregnancy Tumor. Pregnancy tumor or granuloma
gravidarum is a reactive lesion with the same clinical features
to pyogenic granuloma. The lesion may emerge during the
first trimester with a gradual increasing incidence to the
seventh months of pregnancy, which is presumably related
to the rising levels of estrogen and progesterone. Some of
these lesions resolve spontaneously after delivery or undergo
fibrous maturation mimicking an irritation fibroma [4, 10].
The lesions do not occur in people with optimum oral
hygiene suggesting local irritation as an important etiologic
factor [4]. Patients with small isolated lesions and otherwise
healthy gingivae might be monitored for shrinkage after
delivery, but large lesions or generalized pregnancy gingivitis
or periodontitis warrants the need for treatment during
pregnancy [4].

3.1.3. Irritation Fibroma. Irritation fibroma or focal fibrous
hyperplasia is the most common tumor-like lesion of the
oral cavity with the prevalence of 1-2% in general population.
It appears as an asymptomatic, pedunculated, or sessile
exophytic lesion with a smooth surface and being similar to
the surrounding mucosa in color (Figure 5). However, the
surface may show hyperkeratosis from secondary trauma. It
can be firm and resilient or soft with spongy consistency.
Although fibromas usually reach to 1.5 cm or less in diameter
they might appear as very tiny to quite large lesions. The most
commonly affected site is the buccal mucosa along the line of
occlusion; however it can occur anywhere in the oral cavity.
The labial mucosa, tongue, and gingivae can be involved
as well. It is likely that many fibromas represent fibrous
maturation of a preexisting pyogenic granuloma. There is a
female predilection with female to male ratio of 2:1 with
the majority of cases being reported in the fourth to sixth
decades of life. Conservative surgical excision is the treatment
of choice for irritation fibroma with a low recurrence rate.

3.1.4. Peripheral Ossifying Fibroma. Peripheral ossifying
fibroma also called peripheral fibroma with calcification,
ossifying fibroid epulis, and calcifying fibroblastic granuloma
is a reactive gingival enlargement [10]. It represents 2% to 9%
of all gingival lesions and 3% of all oral lesion biopsy samples
[12, 13]. The lesion appears as a smooth, pedunculated, or
sessile, firm to hard mass that usually emanates from the
interdental papilla. It is a red to pink lesion often less than
2 cm in diameter, but lesions up to 8 cm have been reported
as well [10, 13]. Tooth mobility, tooth migration, and bone
destruction have been noticed in some cases [13]. This lesion
occurs exclusively on the gingivae with up to 60% of cases
being reported in the anterior areas of the maxilla (incisor-
cuspid region) [12, 13]. Two theories have been proposed
to explain the pathogenesis of the lesion: it might originate
from a calcified pyogenic granuloma, or it may arise from
an overgrowth and proliferation of different components of
connective tissue in the periodontium, but the main etiology
is yet to be elucidated [12, 14]. Peripheral ossifying fibroma
is predominantly a lesion of teenagers and young adults with a
peak prevalence being between 10 to 19 years [10]. Females are
more affected than males, mainly during their second decade
of life, due to fluctuations of estrogen and progesterone [12].
Treatment usually involves surgical excision, and the lesion
should be excised down to the periosteum. The recurrence
rate has been estimated to be between 8% and 20% [10, 12].

3.1.5. Peripheral Giant Cell Granuloma. Peripheral giant cell
granuloma is a common tumor-like lesion of the oral cav-
ity that conveys a reactive response in the periodontium,
periodontal ligament and gingivae. It occurs exclusively on
the edentulous alveolar ridge and gingivae as a smooth,
reddish-blue, pedunculated, sessile, or nodular mass, which
is firm to palpitation. In some cases the clinical appearance
of the lesion is similar to pyogenic granuloma; however
peripheral giant cell granuloma is more bluish-purple colored
Figure 6: PGCG with an ulcerated, smooth surface and purplish color located buccolingually on the left mandibular ridge.

Figure 7: Massive epulis fissuratum presenting as an exophytic lesion with smooth surface associated with an ill-fit mandibular denture.

as compared with bright red color of pyogenic granuloma (Figure 6) [10, 15]. It is usually less than 2 cm in diameter, but larger sizes are seen occasionally. Progressive growth in some cases may lead to bone and root resorption [16]. The mandible is more affected than the maxilla, and there is a female predilection with female to male ratio of 2:1 [15, 16]. As the giant cells are found to act as a potential target for estrogen, it is not surprising that the lesions are triggered by sex hormones [17]. The lesions can develop at any age; however peak prevalence was found in the fifth and sixth decades of life [10]. Hyperparathyroidism should be considered in differential diagnosis in case of multiple lesions especially with a history of recurrences. In spite of adequate treatment, children with hypophosphatemic rickets are also at a higher risk for developing such lesions [16, 17]. Complete surgical excision with elimination of the entire base of the lesion is the accepted treatment plan for this lesion [15–17].

3.1.6. Epulis Fissuratum. Epulis fissuratum or denture-induced hyperplasia is a reactive lesion of the oral cavity caused by low grade chronic trauma from dentures [18]. About 70% of patients wear ill-fit dentures continuously all day long for more than 10 years [19]. The lesion appears as an asymptomatic single fold or multiple folds of hyperplastic tissues in the alveolar vestibule along denture flanges with a smooth surface, soft to firm consistency, and a normal coloration (Figure 7) [18, 20]. In some cases, severe inflammation or ulceration may be seen in the bottom of the folds [18]. It has been reported in 5% to 10% of the jaws fitted with dentures and is more prevalent in the maxilla than the mandible, especially on the facial aspect of the alveolar ridges [18, 19]. The anterior portions of the jaws are more often affected; however epulis fissuratum of the soft palate has been reported in the literature [20]. Two-thirds to three-fourths of all cases have been found in females [19, 20]. In women, postmenopausal hormonal imbalance makes the oral mucosa susceptible to a hyperplastic growth [18]. It is more frequent in patients over 40 years; however this entity has been reported in patients from childhood to elderly [1].

The size of the lesion varies from less than 1 cm in diameter to massive lesions involving extensive areas on the vestibule [19, 20]. Epulis fissuratum can be treated conservatively or surgically depending on the size of the lesion.

3.1.7. Leaf-Like Fibroma. Ill-fit dentures worn for many years can cause benign hyperplastic fibrous growths or epulides. When a fibrous epulis forms underneath the palatal base of a denture, it is known as a leaf-like denture fibroma or fibroepithelial polyp [21]. It is characterized as a painless, flattened, pink, firm mass attached to the palate by a narrow stalk or a broad base (pedunculated or sessile) which can reach to several centimeters in diameter. The edge of the lesion is serrated resembling a leaf. In most cases, the flattened mass is closely located to the palate and sits in a slightly cupped-out depression [10, 22]. Treatment is accomplished by means of conservative surgical removal and fabrication of new dentures. While altering the denture may decrease the size of the lesion, adjustment alone will not lead to complete regression due to dense nature of the scar tissue. Recurrence is not uncommon [21].

3.1.8. Epulis Granulomatous. Epulis granulomatous or epulis hemangiomatosis—a variant of pyogenic granuloma—presents as an overgrowth arising from a recently extracted tooth socket [23–25]. The precipitating factor in most patients is sharp specula of the alveolar bone left in the walls of the socket [1]. Clinically, it is characterized by reddish, smooth, pedunculated or sessile, nontender, rapidly growing mass with soft to firm in palpation. Surface of the lesion may be ulcerative due to secondary trauma [23, 24]. It often bleeds easily because of its high vascular content [23]. The growth may become apparent in one or two weeks after tooth extraction [1]. Lesions of larger sizes were also incidentally seen as an oral finding in patients with Klippel-Trenaunay syndrome [23]. Evaluation of socket and removal of any bony spicules or tooth fragments at the time of extraction prevent formation of an epulis granulomatous. Excision of the raised mass and curettage of the alveolus to ensure the
elimination of irritating particles is needed to treat the lesion [23–25].

3.1.9. Pulp Polyp. Pulp polyp or chronic hyperplastic pulpitis or pulpitis aperta is an uncommon reactive lesion, which occurs when caries have destroyed the tooth crown [1]. It appears as a smooth, soft to firm, red to pink, pedunculated, or sessile mass occupying the entire carious cavity in the affected tooth resembling an enlarged gingival tissue (Figure 8) [26, 27]. The size of the lesion varies from less than 1 cm in diameter to large masses (about 4 cm) [26, 27]. It is most frequently found in the deciduous and permanent first molars of children and young adults and is a rare phenomenon in the middle-aged adults [1, 28]. It is usually asymptomatic, but discomfort can occur during mastication. Response to electrical and thermal stimuli may be normal [28]. Periapical radiographs may show an incipient chronic apical periodontitis when pulp involvement is extensive or lingering [28]. The polyp may cover most of the remaining crown of the tooth, giving the lesion an appearance of a flashy mass [29]. A similar hyperplastic mass around a draining sinus tract of a tooth with pulpoperiapical pathology is called parulis [4]. The treatment plan of this lesion includes endodontic treatment or tooth extraction [28].

3.1.10. Giant Cell Fibroma. Giant cell fibroma is a fibrous hyperplastic soft tissue lesion classified as an inflammatory hyperplasia [1]. However, a controversy exists about the origin and etiology of this entity [30]. It represents approximately 2% to 5% of all oral fibrous lesions and 0.4% to 1% of all oral biopsy samples [31]. Clinically, it appears as an asymptomatic, sessile, or pedunculated mass with rough surface (papillary or granular) and firm in palpation (Figure 9) [30, 31]. Despite previously mentioned reactive lesions, giant cell fibroma did not have a completely smooth surface; hence it may clinically be mistaken with lesions of squamous epithelium origin such as papilloma [10]. The color of the lesion is pink or similar to the surrounding normal mucosa and is usually less than 1 cm in diameter [10, 31]. In about 60% of cases, the lesion is diagnosed in the second to third decade of life with only 4% to 17% of giant cell fibromas being found in children younger than 10 years [31, 32]. There is a slight female predilection and gingivae are the most affected site (about 50% of cases) followed by the tongue, palate, buccal mucosa, lips, and floor of the mouth [10, 31]. Moreover, mandibular gingivae are affected twice as often as the maxillary gingivae [32]. Giant cell fibroma usually is treated by conservative surgical excision. Electrosurgery and laser therapy have been suggested as alternative modalities especially in children. Recurrence is rare [10, 28].

3.1.11. Inflammatory Papillary Hyperplasia. Inflammatory papillary hyperplasia (palatal papillomatosis and denture papillomatosis) is a reactive lesion seen most often in patients with an ill-fit denture wearing all-day-long and poor oral hygiene [10, 33]. This condition is encountered in 10% to 20% of denture wearers [1, 10]. It is featured by exophytic masses with pebbly or cobblestone appearance on the hard palate beneath a denture base with or without symptoms [10]. Although these lesions appear as red, soft masses in the inflammatory stage they convert to pink and firm when they mature to fibrous stage [1]. In some cases, denture papillomatosis develops on the edentulous mandibular alveolar ridge or on the surface of an epulis fissuratum [10]. It can occur at any age. However, it is most frequently encountered in the third to fifth decade of life with a male predilection [33]. In addition to frictional irritation provoked by loose-fit dentures, Candida albicans has an etiologic role in this entity [1]. On rare occasions, this condition occurs on the palate of patients without denture especially in people who habitually breathe through their mouth or have a high palatal vault [10]. Less extensive lesions are resolved by removing the denture at night and improving oral hygiene. Patients can also benefit from antifungal agents. Various surgical methods have been suggested for the treatment of this lesion such as partial thickness or full thickness surgical blade excision, curettage, electro surgery, and cryosurgery [10, 34, 35].
3.2. Salivary Gland Lesions. A wide range of lesions arise from intraoral salivary glands, which are categorized as nonneoplastic and neoplastic entities. According to a 15-year retrospective study by Mohan et al., 55% of lesions were nonneoplastic and 45% were neoplastic [36]. When salivary gland lesions appear as an exophytic lesion they are usually characterized by a dome-shaped or nodular mass with a smooth surface and fluctuant to firm in palpation [1]. The most common lesions of salivary gland origin are as follows (Table 2).

3.2.1. Mucocele and Ranula. Mucocele is a common nonneoplastic lesion of salivary gland origin resulting from breaking or dilation of salivary ducts secondary to obstruction or local trauma [37]. Clinically, it appears as an asymptomatic, fluctuant to firm, bluish to pink, nodular, or dome-shaped mass with a smooth surface (Figure 10) [1,37]. The size of the lesion varies from 1-2 mm to several centimeters but remains smaller than 1.5 cm in diameter [37]. The incidence is pretty high, 2.5 cases per 1000 persons with no sex predilection [38]. Lesions last from a few days to several months. Many patients mention a history of recurrent swelling with periodic rupture and release of fluid content [10, 38]. The majority of cases have been reported in the first three decades of life, and it is rare among children younger than 1 year of age [37, 38]. The lower lip is the most frequently (60%) affected site followed by buccal mucosa, anterior ventral tongue, floor of the mouth, and upper lip [10]. In some cases, mucoceles rupture and heal spontaneously; however conventional surgical treatment may be suggested in large lesions. Some alternative modalities such as cryosurgery, intralesional injection of steroids, and laser therapy have been implemented as well [37].

When a mucocele occurs in the floor of the mouth it is called ranula featuring as a bluish, dome-shaped or nodular; fluctuant exophytic lesion with a smooth surface usually located lateral to the midline [10]. Fluctuation in size considered as a pertinent feature noticed in the history of ranula. The lesion measures its smallest size early in the morning and reaches to the largest scale at the time of meals [1]. A prevalence of 0.2 cases per 1000 persons has been found, and it accounts for 6% of all oral sialocysts [39]. Most of the patients are young adults with peak frequency being in the second decade of life [39]. This lesion is treated by marsupialization or removal of the feeding sublingual gland; however marsupialization leads to recurrence in 25% of cases [1, 10].

3.2.2. Pleomorphic Adenoma. Pleomorphic adenoma or mixed tumor is a benign salivary gland tumor which constitutes 3% to 10% of head and neck neoplasms and about 1% of all body tumors [40, 41]. It is the most common (73%) tumor of both minor and major salivary glands [41, 42]. Corresponding to minor salivary glands, the palate is the most affected site followed by lips, buccal mucosa, tongue, floor of the mouth, pharynx, retromolar area, and the nasal cavity [41]. It is suspected when a clinician encounters a unilateral, asymptomatic, slow growing, firm, nodular, or dome-shaped mass covered by a normal-colored mucosa (Figure 11) [40, 41]. The surface of the lesion is usually intact, but ulceration of the overlying mucosa has been reported in some instances [40]. The size of the lesion varies from 1 to 5 cm with the average of 2.6 cm in diameter [41]. In large cases bone resorption has been reported [40]. It can occur at any age, but majority of patients have been reported in their fifth to sixth decades of life [41, 42]. Lip lesions tend to occur at an earlier age than that of other locations [41]. There is a female predilection with female to male ratio of 7:3 [41]. Malignant transformation to pleomorphic adenocarcinoma has been noticed in about 6% of cases [42]. Surgical excision usually yields promising results treated with good prognosis. A recurrence rate of 2% to 44% has been reported, and patients younger than 30 years are more likely to have relapsing lesions [41]. Risk for recurrence seems to be lower in the minor salivary gland tumors than that of major salivary glands [41].

3.2.3. Mucoepidermoid Carcinoma. Mucoepidermoid carcinoma or mucoepidermoid tumor is the most common malignant salivary gland neoplasm. It accounts for about 3% of all head and neck tumors [10, 43]. Two-thirds of the lesions arise within the parotid gland and one-third within the minor salivary glands. Intraorally, palate is the most affected site followed by retromolar area, floor of the mouth, buccal mucosa, lips, and tongue. Intraosseous lesions have been also reported in some cases [10, 43, 44]. The tumor can occur at
| Entity                        | Age                | Gender | Site of involvement | Surface texture | Type of base           | Consistency       | Color           | Size      | Symptom & sign                                      | Treatment                                 | Recurrence |
|-------------------------------|--------------------|--------|---------------------|-----------------|------------------------|------------------|-----------------|-----------|----------------------------------------------------|-------------------------------------------|------------|
| Mucocele & ranula             | First 3 decades of life | No sex predilection | Lower lip/floor of the mouth | Smooth          | Nodular/dome shaped    | Fluctuant to firm | Blush to pink | <1.5 cm  | Asymptomatic                                       | Marsupialization                         | 25%        |
| Pleomorphic adenoma           | 5-6th decades      | Female | Palate              | Smooth          | Nodular/dome shaped    | Firm             | Normal colored | 1-5 cm   | Asymptomatic                                       | Surgical excision                         | 2-44%      |
| Mucoepidermoid carcinoma      | 3-6th decades      | Female | Palate              | Smooth          | Nodular/dome shaped    | Firm             | Pink/blush to red or normal colored | Up to several cm | Asymptomatic/Painful in high Grades | Wide surgical excision                     | Up to 60%   |
| Adenoid cystic carcinoma      | 5-6th decades      | Female | Palate              | Smooth/ulcerative | Nodular           | Firm             | Pink/normal colored | Up to several cm | Pain is common/bone destruction/distant metastasis | Local radical excision + radiotherapy/chemotherapy | —          |
any age, but the majority of patients have been diagnosed between the third and sixth decade of life with a female predilection. Although it is rare in children, mucoepidermoid carcinoma is the most common malignant salivary gland neoplasm in childhood [10, 43–45]. Clinically, it appears as a smooth-surfaced nodular or dome-shaped, firm mass with a diameter up to several centimeters. The color of the lesion may be pink, bluish to red, or even similar to the surrounding normal mucosa (Figure 12) [1, 10, 43]. Patients are usually asymptomatic, but high grade tumors might be painful [44]. Palatal neoplasms cause bone resorption, and an ulcerative surface might be found in some tumors. [10, 44]. Low grade neoplasms are treated by surgical excision with free surgical margins. High grade ones require wide surgical excision, neck dissection, and postoperative radiotherapy. Recurrence rate is up to 60%, which mostly occurs within 1 year after treatment [45].

3.2.4. Adenoid Cystic Carcinoma. Adenoid cystic carcinoma is the most common malignant tumor originated from the submandibular and minor salivary glands. It comprises 5% to 10% of all salivary gland neoplasms and 2% to 4% of all head and neck malignancies [4, 46]. It can occur in any salivary gland, but approximately 50% to 60% of cases develop within minor salivary glands. The palate is the most frequently affected site in the oral cavity. Clinically, it is characterized by a slow growing, pink to normal-colored nodular mass with a smooth surface and firm consistency [1, 46, 47]. An intraoral adenoid cystic carcinoma may exhibit mucosal ulceration, a feature that helps distinguish it from a benign pleomorphic adenoma [4]. Pain is a common finding and usually occurs early in the course of the disease before an apparent swelling develops [10]. In cases arising in the palate or maxilla, there is an evidence of bone destruction adjacent to the tumor [4, 47]. Local recurrences, perineural spreading, and distant metastases have been also reported as important features of this entity [4]. The peak prevalence of this tumor is in the fifth and sixth decades, and it is rare in people younger than 20 years [10, 46]. An almost equal sex distribution has been mentioned; however some studies showed a female predilection [10, 46, 47]. Because of the ability of this lesion to spread along the nerve sheets, radical surgical excision is suggested as the accepted treatment plan. The tumors of the minor salivary glands should be treated by local radical excision and postoperative radiotherapy. Chemotherapy is also recommended in the management of advanced and metastatic salivary gland tumors [47].

3.3. Mesenchymal Lesions. Peripheral oral mesenchymal tumors are considered among uncommon lesions of the oral cavity (Table 3). According to a 10-year retrospective study by Mendez, only 4% of oral lesions were of mesenchymal origin [48]. Clinically, these lesions present as asymptomatic, slow growing, nodular, or dome-shaped masses with a smooth surface and firm consistency. These lesions are usually covered with normal mucosa unless chronically traumatized. They can be involved any site of the oral cavity [1]. The most common oral lesions with mesenchymal origin are lipoma, neurofibroma, schwannoma, lymphoma, hemangioma, and lymphangioma [49].

3.3.1. Neurofibroma. Neurofibroma is the most common peripheral nerve tumor; however it is rarely seen in the oral cavity. Most cases of oral neurofibroma are multiple and as a part of neurofibromatosis syndrome, but it rarely appears as a solitary mass without visceral manifestations [10, 50]. It has been demonstrated that solitary neurofibroma is a hyperplastic hamartomatous malformation rather than a neoplastic lesion [51]. It appears as an asymptomatic, slow growing, soft to firm, nodular, or sessile mass with a smooth surface (sometimes lobulated) and pink in coloration (Figure 13) [50, 51]. It most commonly develops on the tongue followed by palate, mandibular ridge/vestibule, maxillary ridge/vestibule, buccal mucosa, lips, floor of the mouth, and gingivae with up to several centimeters in diameter [50, 51]. Intraosseous lesions have been also reported in the posterior mandible as a well-defined or poorly defined unilocular or multilocular radiolucency [51, 52]. It is most commonly observed in young adults with a peak prevalence in the third decade of life with the sex predilection being still debatable [50, 51]. The tumor is usually treated by complete excision.
Table 3: General characteristics of smooth-surfaced oral exophytic lesions of mesenchymal origin.

| Entity       | Age       | Gender       | Site of involvement | Surface texture | Type of base | Consistency | Color         | Size          | Symptom & sign                                                                 | Treatment                                                                                           | Recurrence |
|--------------|-----------|--------------|---------------------|-----------------|--------------|-------------|---------------|---------------|--------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------|------------|
| Neurofibroma | 3rd decade| No sex predilection | Tongue             | Smooth          | Nodular/sessile | Soft to firm | Pink          | Up to several centimeters | Asymptomatic                                          | Complete excision                               | Rare        |
| Schwannoma   | Average: 34 years | Female | Tongue             | Smooth          | Nodular/sessile | Rubbery      | Same to normal mucosa | 0.5–4 cm | Asymptomatic                                            | Surgical excision                                      | Rare        |
| Lipoma       | >40 years | Male         | Buccal mucosa/vermilion | Smooth          | Nodular/sessile/ pedunculated | Soft/fluctuant | Pink to yellowish | <3 cm (often) | Asymptomatic                                                | Conservative surgical excision                        | High in intramuscular lesions                   |
| Lymphoma     | Average: 59 years | Male | Buccal vestibule & postpalate | Smooth/ulcerative | Nodular | Soft to firm | Pink/purplish/normal | Up to several cm | Non-tender painful in intraosseous lesions | Chemotherapy/radiotherapy/Sclerotherapy/ Surgical excision/laser therapy/cryotherapy | —          |
| Hemangioma   | Early infancy | Female | Lips              | Smooth/fibulated | Sessile | Soft          | Pink to red purple | Up to several cm | Asymptomatic                                                | —                                                                                      | —          |
| Lymphangioma | 1st decade | No sex predilection | Tongue             | Smooth/pebbly   | Sessile | Soft          | Pink to yellowish or normal color | Up to several cm | Macroglossy/airway obstruction/sialorrhea/jaw deformity | —                                                                                      | —          |
with a low recurrence rate. However, neurofibroma may convert to neurofibrosarcoma in 5% to 15% of cases especially in multiple lesions [51].

3.3.2. Schwannoma. Schwannoma, also called neurilem-moma, neurinoma, or perineural fibroblastoma, is a benign tumor of neuroectodermal origin [53, 54]. Approximately 25% to 45% of lesions are seen in the head and neck with 1% being reported in the oral cavity as well [53]. Clinically, it is characterized by solitary, asymptomatic, rubbery, nodular, or sessile mass with a smooth surface and is similar to the normal mucosa in coloration (Figure 14). The size of the lesion varies from 0.5 to 4 cm in diameter [53, 54]. Intraoral lesions are frequently located in the tongue followed by palate, floor of the mouth, buccal mucosa, gingivae, lips, and vestibular mucosa [55]. Schwannoma can occur as an intraosseous lesion, which accounts for 1% of all benign primary bony tumors [54, 55]. Intraosseous lesions appear as either unilocular or multilocular radiolucencies [53]. Peripheral lesions are usually painless, but tenderness may occur in some instances [10]. Pain and paresthesia are not uncommon for intrabony tumors [10]. Schwannoma has been reported in the age range of 8 to 72 years with an average age of 34 years with a slight female predilection (female to male ratio of 1.6:1) [53, 55]. Surgical excision is the treatment of choice and recurrence and malignant transformation are extremely rare [55].

3.3.3. Lipoma. Lipoma is a benign tumor, which seldom occurs in the mouth. It constitutes 4% to 5% of all benign tumors in the body representing about 1% to 5% of all neoplasms of the oral cavity [56]. Generally, the prevalence of the lesion is balanced in both genders; however a slight male predilection has been reported. Lipoma is a rare entity in children, and most of the patients are over 40 years [57]. It presents as a slow growing, nodular, sessile, or even pedunculated mass with a smooth surface and fluctuant to soft in palpation. The superficial lesions usually show yellowish hue, while more deeply seated ones appear as a pink mass [10, 56, 57]. Lipoma varies in size from small to large masses mostly measuring less than 3 cm in diameter [10, 56]. The buccal mucosa and buccal vestibule are the most commonly affected sites, which accounts for about 50% of all cases [10]. Less affected sites include the tongue, floor of the mouth, and lips [10]. Lipoma is treated by conservative surgical excision and usually does not recur. Intramuscular lipoma has a somewhat higher recurrence rate because of problems to remove completely [4].

3.3.4. Lymphoma. Lymphomas are heterogeneous malignant neoplasms of the lymphocyte cell lines. They can be divided as Hodgkin and non-Hodgkin lymphoma (HL and NHL). Hodgkin lymphoma rarely shows extra-nodal disease (1% of cases) unlike NHL, which arises from extra-nodal sites in 20% to 30% of patients [58, 59]. Extra-nodal lymphoma is the second most commonly encountered neoplasm after squamous cell carcinoma in the head and neck region, which accounts for 5% of all malignancies of head and neck [59–61]. Oral lymphomas usually occur secondary to a more widespread involvement through the body; however it can present as a primary lesion in the oral cavity with the prevalence of 0.1% to 2% [60]. Clinically, it appears as a nontender, smooth surface, soft to firm mass in the mouth (Figure 15) [10, 58–60]. The size of lesion varies from small to large with a pink, purplish, or normal coloration. The surface of lesion may be intact or ulcerative [10, 59, 60]. The mean age of patients with lymphoma is 56 years, and there is a male predilection [60, 61]. The most affected site for oral lesions is buccal vestibule, posterior palate, and gingivae [10, 60]. Intraosseous lesions were also reported in some cases. Bony lesions often present with low grade pain, which can mimic toothache [60]. Radiotherapy plus chemotherapy is recommended for intermediate and high grade tumors. A failure rate of 30% to 50% was demonstrated in intermediate grade lesions; however high grade lesions show a 60% mortality rate [10].

3.3.5. Hemangioma. Hemangioma is a benign common tumor in the head and neck region, but relatively rare in the mouth. In the oral cavity, it can cause esthetic and functional impairment depending on its location and size [62]. The peak prevalence of this entity is described soon after birth or in early infancy; however some cases have been reported
in adulthood [10, 62]. It is noted that hemangioma is the most common tumor of infancy, occurring in 5% to 10% of one-year-old children [10]. Oral lesions are most common in the lips, gingivae, tongue, and buccal mucosa [62, 63]. Eighty percent of cases occur as a single lesion, but 20% of affected patients present multiple tumors [10]. Clinically, it is characterized by an asymptomatic, soft, smooth, or lobulated, sessile mass with various sizes from a few millimeters to several centimeters (Figure 16). The color of the lesions ranges from pink to red purple, and tumor blanches on the application of pressure [63, 64]. Gingival lesions which arise from the interdental papillae can spread laterally to involve adjacent teeth [63]. There is a female predilection with female to male ratio of 3:1, and they usually occur in whites more often than other racial groups [10]. Treatment of hemangioma depends on its size and location. Small lesions are treated by sclerotherapy, conventional surgical excision, laser therapy, and cryotherapy. In large cases, treatment should include embolization or obliteration of the lesion and the adjacent vessels [62].

3.3.6. Lymphangioma. Lymphangioma is a benign, hamartomatous tumor of lymphatic vessel origin. It has a marked tendency for the head and neck region in a way that 75% of all cases occur in this area. Almost half of the lesions are noted at birth and about 90% developed by two years of age [65]. In the oral cavity, lymphangioma mostly occurs on the dorsal surface and lateral borders of the tongue followed by gingivae, buccal mucosa, and lips [65, 66]. Pathognomonic features of tongue lymphangioma are irregular nodularity of the surface with gray to pink projections and macroGLOSSIA [66]. There is no sex predilection, and oral lesions are most common in the first decade of life [66]. The clinical manifestation of the lesion varies based on whether it is superficial or deep. Superficial lesions appear as soft exophytic masses with rough (papillary or pebbly) surface and pink to yellowish coloration. Deeper lesions are described as soft diffuse masses with normal color and smooth surface [65, 67]. Massive lesions might cause macroGLOSSIA, obstruction in upper air way, sialorrhea, and jaw deformity as well as difficulties in mastication, speech, and oral hygiene [65]. Lymphangioma is treated by surgical excision, cryotherapy, electrocautery, sclerotherapy, steroid administration, embolization, laser surgery, and radiation therapy [65].

3.4. Lesions with Rough Surface. This group of lesions most frequently occurs as a result of epithelial proliferation due to reaction to human papilloma virus or a neoplastic process (Table 4).

3.4.1. Squamous Papilloma. Oral squamous papilloma is a benign proliferation of the stratified squamous epithelium. It occurs in one of every 250 adults and constitutes approximately 3% of all oral biopsy lesions [10, 68]. The etiologic factor is the human papillomavirus (HPV), and viral subtypes 2 and 11 have been isolated from up to 50% of oral papillomas [10]. Clinically, it appears as a single, asymptomatic, pedunculated or sessile, soft to firm mass with verrucous, granular, or papillomatous surface (Figure 17). The lesion may be white, slightly red, or normal in color [4, 68–70]. Papilloma usually enlarges rapidly to a maximum size of 5 mm with little or no change in diameter thereafter [10]. The most affected sites in the oral cavity are palatal mucosa and the tongue, but any oral surface may be involved [69]. Although papilloma can involve patients at any age it is diagnosed most often in people between 30 and 50 years [10]. Surgical excision by either scalpel or laser ablation is the treatment of choice. However, other modalities such as electro cautery, cryosurgery, and intralesional injection of interferon have been suggested [70]. Recurrence is uncommon, except for patients infected with human immunodeficiency virus [70].

3.4.2. Verruca Vulgaris. Verruca vulgaris is a benign virus-induced hyperplasia of stratified squamous epithelium. It is usually encountered in children with a peak incidence between 12 and 16 years. However, about 10% of general population affect the disease in their middle age [71, 72]. Most of oral lesions are located on the vermilion border, labial mucosa, or anterior tongue [4, 10]. Clinically, it appears as an exophytic, sessile, or pedunculated lesions with papillary projections (verrucous or papillomatous) or a rough pebbly surface [10, 71]. The lesions are asymptomatic and may be pink, yellowish, or white in coloration [10]. It usually
| Entity                        | Age            | Gender       | Site of involvement          | Surface texture       | Type of base            | Consistency | Color               | Size          | Symptom & sign     | Treatment                                                                 | Recurrence                        |
|------------------------------|----------------|--------------|------------------------------|-----------------------|-------------------------|-------------|---------------------|---------------|-------------------|---------------------------------------------------------------------------|-----------------------------------|
| Squamous papilloma           | 30–50 years    | No sex predilection | Palate & tongue              | Verrucous/granular/papillomatous | Pedunculated/sessile    | Soft to firm | White/slightly red/normal | Maximum 5 mm | —                | Surgical excision/electrocautery/cryosurgery/intralesional interferon | Uncommon except for patients with HIV infection |
| Verruca vulgaris             | 12–16 years    | —            | Vermilion border/labial mucosa | Verrucous/papillomatous/Pebble | Sessile/pedunculated    | Soft to firm | Pink/yellowish/white | Few mm        | Asymptomatic | Conservative surgical excision | Low                              |
| Verrucous carcinoma          | >6th decade    | Male         | Vestibular mucosa            | Verrucous             | —                       | Firm        | White to normal      | Up to several cm | Asymptomatic | Wide surgical excision/radiotherapy/chemotherapy | —                                |
| Squamous cell carcinoma      | 62 years       | Male         | Tongue                       | Verrucous             | —                       | Firm to hard | White/pink/red       | Up to several cm | Painless/moth-eaten radiolucenty | Wide surgical excision/radiotherapy/Surgery/laser/cryosurgery/electrosurgery | 3.3%                             |
| Multifocal epithelial hyperplasia | Childhood   | Female       | Labial and buccal mucosa     | Cobblestone           | —                       | Soft to firm | White to normal      | Up to several cm | Nontender |                                                                                                     | Minimum                           |
enlarges rapidly to its maximum size and remains constant for months or years thereafter [71, 72]. Approximately 23% of lesions show spontaneous regression in two months and the remaining lesions within 2 years [72]. Oral lesions are usually treated by conservative surgical excision. Meanwhile, laser therapy, cryotherapy, or electrosurgery has been also recommended. A small proportion of treated lesions show recurrence [4, 10].

3.4.3. Verrucous Carcinoma. Verrucous carcinoma which also called Ackerman tumor, Buschke-Lowenstein tumor, florid oral papillomatosis, epithelioma cuniculatum, or snuff dipper’s cancer is a nonmetastasizing low grade variant of oral squamous cell carcinoma [68, 73]. The incidence rate has been estimated as one lesion per 1,000,000 of the population each year. The etiology of the lesion is not clear, but oral lesions predominantly occur in patients habituated to areca chewing, alcohol drinking, smoking, and having poor oral hygiene. However, 15% to 51% of the lesions have been found in people without these habits [10, 68]. The most common affected sites in the oral cavity are vestibular mucosa, buccal mucosa, gingivae, and the tongue [10, 68, 73]. There is a male predilection with the majority of lesions being reported in the sixth decade [4, 68, 73]. The lesion is featured by an asymptomatic, broad-based, well-circumscribed, thick, pink to white plaque resembling a cauliflower with verruciform surface (Figure 18) [10]. Similarly simultaneous oral and genital lesions of larger sizes would suggest condyloma acuminatum [10]. Malignant transformation has been detected in 20% of the lesions [73]. Wide surgical excision, radiotherapy, and chemotherapy have been recommended for treatment.

3.4.4. Squamous Cell Carcinoma. Squamous cell carcinoma (SCC) accounts for 90% of all oral cancers and is considered as the eighth most frequent cancer globally [74]. Males are affected more frequently than females with a male to female ratio of 3:1 [10]. The median age of diagnosis is 62 years. However, the incidence of oral SCC in persons younger than 45 is increasing [75–77]. Tongue is the mostly affected site in the oral cavity followed by floor of the mouth, gingivae, palate, retromolar area, buccal, and labial mucosa [10, 75–78]. In young patients and those with congenital oral squamous cell carcinoma the tongue is the mostly affected site of involvement as well [10]. Warning signs of oral cancer include red-white lesions, ulcer lasting more than three weeks, pain of the tongue due to its mobility and sensitive nature, lump in the oral cavity or in the neck area, discomfort with speech or swallowing, mobile teeth in the absence of periodontitis, anesthesia, and earache without apparent disease. The most frequent complaints among oral cancer patients are swelling, pain, and ulceration [74]. Clinically, it can appear in various forms such as a red or white plaque, a solitary chronic ulcer or an exophytic mass [75]. Approximately 55% of the tongue lesions were exophytic-type. This type of lesions has a rough surface (verrucous) and usually is irregular in shape (Figure 19). The lesion has a broad base and it is white, pink, or red in color. Ulceration may be present in larger fungating lesions with necrotic and multicolored surface. In addition, oral SCCs are painless and firm to hard in palpation and bleeding is not an early characteristic feature [1, 10, 75]. However, in cases with destruction of the underlying bone pain may be reported, and a “moth-eaten radiolucency” with ill-defined or ragged borders is found [10]. Despite early oral cancer whose manifestations are not definitive late-stage oral SCC show quite prominent signs such as a large mass with irregular margins, ulceration, nodularity, and fixation to the surrounding tissues [74]. The treatment of intraoral SCC is guided by the clinical stage of the disease and consists of wide surgical excision, radiation therapy, or a combination of surgery and radiation therapy [10]. The recurrence rate is estimated about 33% within 2 to 96 months [79].

3.4.5. Multifocal Epithelial Hyperplasia. Multifocal epithelial hyperplasia, also known as focal epithelial hyperplasia, multifocal epithelial papilloma, virus epithelial hyperplasia, and Heck disease, is a virus-induced proliferation of oral squamous epithelium [10]. Although this entity is usually a childhood condition other age groups may also be affected. The frequency of the lesion ranges from 0.002 to 35% in different populations and 70% of patients are in first two decades of their life [79]. In addition, there is a female predilection with female to male ratio of 3:1 [79]. It appears
as multiple, soft, circumscribed, nontender, flattened, or rounded papules, with either whitish color or color similar to the normal mucosa and cobblestone appearance (Figure 20) [10, 79, 80]. Most of the lesions are less than 0.5 cm, but lesions with several centimeters in size have been reported as well [10, 79]. The most frequent affected sites are labial, buccal, and lingual mucosa followed by gingivae, palate, and tonsillar mucosa [10]. There is no potential for malignant transformation [79, 80]. However, Niebrügge et al. showed malignant transformation in a female patient with long-standing Heck disease [81]. It is a self-limiting disease; hence it is recommended that only lesions located in areas subjected to trauma be excised. Risk of recurrence after therapy is minimal. There are several treatment modalities like scalpel surgery, cryosurgery, electrosurgery, and carbon dioxide laser vaporization [10, 79].

4. Discussion

In this review article we proposed a practical diagnostic decision tree for oral peripheral exophytic lesions as well as a brief overview about each entity. Although this group of lesions comprises a constellation of heterogeneous origins and pathogeneses, some clinical features help us categorize them to come arrive a more timely and precise differential diagnosis. We used surface texture for clinical classification of the lesions into two major groups: “lesions with smooth surface” and “lesions with rough surface.” Furthermore, the former was divided into three subgroups constituting 23 entities, and the latter constituting six lesions. While general characteristics guide to a certain group of lesions, special features point to a unique entity.

In regard to reactive hyperplastic lesions, detecting the insult factor through history taking or physical examination plays the key role for definite diagnosis. For example, a bright red color in a soft and easily bleeding mass suggests a pyogenic granuloma, which is called a pregnancy tumor in a pregnant woman. However a peripheral giant cell granuloma presents as a more bluish, buccolingually located mass exclusively on the gingivae or alveolar ridge with a firm consistency. Meanwhile, it has a potential to cause root and bone resorption. Similarly, peripheral ossifying fibroma has a tendency to induce tooth mobility and bone destruction, but it is firm to hard in palpation with a pale pink coloration [4, 9, 10, 12, 13, 15].

Moreover, some lesions can be identified primarily by their region such as those relating to ill-fit dentures. Epulis fissuratum is seen along denture flanges as single or multiple folds, inflammatory papillary hyperplasia is located mostly on the hard palate with a pebbly or cobblestone appearance, and leaf-like fibroma presents with a narrow stalk and serrated borders beneath the base of maxillary dentures [10, 18–21].

Some exophytic lesions are found in close approximation to a tooth such as pulp polyp proliferating inside a large tooth cavity, epulis granulomatous emanating from an extraction socket, and parulis on the alveolar mucosa or attached gingivae of a necrotic tooth [23, 25, 28].

Oral exophytic lesions of nonneoplastic salivary origin such as mucocele and ranula appear as soft and fluctuant dome-shaped lesions on oral mucosa containing salivary minor glands with a history of periodic rupture and release of fluid content. Therefore, they should not be suspected when a lesion is located on the hard palate or gingivae due to lack of submucosal salivary glands. On the other hand, a dome-shaped or nodular lesion on the posterolateral portion of the hard palate with a soft to firm consistency and slow growth rate would prompt the clinician to consider a benign salivary neoplasm such as pleomorphic adenoma in higher rankings of differential diagnosis. Moreover, accompanying pain raises the possibility of a malignant counterpart like adenoid cystic carcinoma or mucoepidermoid tumor. While a dome-shaped lesion on the lower lip is usually suggestive of a mucocele, such lesion in the upper lip should be suspected as a salivary neoplasm [37, 41, 43, 45, 46].

Peripheral oral exophytic lesions of mesenchymal origin are quite infrequent as compared with reactive or salivary lesions. Although they might show similar clinical features special characteristics such as location help differentiate them.

Neoplasms of neuronal origin mostly appear on the tongue as asymptomatic, soft to firm masses with neurofibroma being mostly multiple contrary to solitary schwan- noma. In addition, concomitant eye and skin manifestations indicate neurofibromatosis. A soft fluctuant mass with a yellowish hue especially in the buccal mucosa points out a tumor of adipose tissue (lipoma). Non-Hodgkin lymphoma in the oral cavity produces a nodular pink or purplish mass with an intact or ulcerated surface on the buccal vestibule or palate with a potential for progressive growth and local destruction. Hemangioma and lymphangioma most frequently happen on the tongue sometimes with a rough surface despite their connective tissue origin. Hemangioma is characterized by a pink or purplish color and blanching on pressure, whereas lymphangioma causes macroGLOSSIA and gray to pink projections [1, 4, 10].
As a general rule, lesions originating from the epithelium present with a rough surface. An exception in this regard is focal epithelial hyperplasia with numerous small flat-end papules scattered on the oral mucosa most commonly in children and adolescence. Small finger-like projections on lesion surface are invariably seen in viral-induced lesions such as squamous papilloma, verruca vulgaris, and condyloma acuminatum. Papilloma tends to occur as a single lesion rarely exceeding 1 cm in diameter. On the other hand, verruca vulgaris and Heck disease mostly appear in multiple forms with the former being coexisted with cutaneous lesions. Coincidence of large verruca vulgaris-form lesions in the genitalia and oral cavity as well as high-risk sexual behavior refer to a condyloma acuminatum.

Squamous cell carcinoma when appears as a broad-based exophytic lesion involves the lateral border of the tongue with a rough nonhomogenous or sometimes ulcerative or necrotic surface and a progressive potential to unlimited growth. On the other hand, its nonmetastasizing counterpart—verrucous carcinoma—is mostly encountered on the vestibular or buccal mucosa in a snuff-dipper patient with a homogenous cauliflower appearance and an association with smokeless tobacco use.

5. Conclusion

We proposed a diagnostic decision tree regarding oral peripheral exophytic lesions divided into lesions with smooth surface and rough-surfaced lesions. Upon confronting a peripheral exophytic mass in the oral cavity a clinician should consider some features such as surface texture, shape of base, color, and consistency in order to categorize the lesion and progress along the decision tree to reach a logistic differential diagnosis.

Conflicts of Interest

The authors declare that there are no conflicts of interest regarding the publication of this paper.

References

[1] N. K. Wood and P. W. Goaz, *Differential Diagnosis of Oral And Maxillofacial Lesions*, Mosby, St. Louis, Mo, USA, 5th Edition edition, 1997.

[2] A. B. R. Santosh, D. Boyd, and K. K. Laxminarayana, "Proposed clinico-pathological classification for oral exophytic lesions," *Journal of Clinical and Diagnostic Research*, vol. 9, no. 9, pp. ZE01–ZE08, 2015.

[3] R. B. Zain, N. Ikeda, I. A. Razak et al., "A national epidemiological survey of oral mucosal lesions in Malaysia," *Community Dentistry and Oral Epidemiology*, vol. 25, no. 5, pp. 377–383, 1997.

[4] M. Glick, *Burket's Oral Medicine*, People's medical publishing house, 12th edition edition, 2015, pp. 147-172, 175-188, 236-237.

[5] A. Bermejo-Fenoll and P. López-Jornet, "Differential diagnosis of exophytic lesions of soft oral tissue," *Medicina Oral, Patología Oral y Cirugía Bucal*, vol. 10, no. 5, pp. 470–471, 2005.

[6] N. G. Nikitakis, "Oral soft tissue lesions: a guide to differential diagnosis part II: surface alterations," *Brazilian Journal of Oral Sciences*, vol. 4, no. 13, pp. 707–715, 2005.

[7] V. Reddy, S. Saxena, S. Saxena, and M. Reddy, "Reactive hyperplastic lesions of the oral cavity: a ten year observational study on North Indian population," *Journal of Clinical and Experimental Dentistry*, vol. 4, no. 3, pp. e136–e140, 2012.

[8] H. Kadeh, S. Saravani, and M. Tajiik, "Reactive hyperplastic lesions of the oral cavity," *Iranian Journal of Otorhinolaryngology*, vol. 27, no. 79, pp. 137–144, 2015.

[9] H. Jafarzadeh, M. Sanatkhani, and N. Mohtasham, "Oral pyogenic granuloma: a review," *Journal of Oral Science*, vol. 48, no. 4, pp. 167–175, 2006.

[10] B. W. Neville, D. D. Damm, C. M. Allen, and J. E. Bouquot, *Oral and Maxillofacial Pathology*, Saunders-Elsevier. China, 3rd edition, 2009.

[11] S. R. Gomes, Q. J. Shakir, P. V. Thaker, and J. K. Tavadia, "Pyogenic granuloma of the gingiva: a misnomer?: a case report and review of literature," *Journal of Indian Society of Periodontology*, vol. 17, no. 4, pp. 514–519, 2013.

[12] M. J. Franco-Barrera, M. G. Zavala-Cerna, R. Fernández-Tamayo, I. Vivanco-Pérez, N. M. Fernández-Tamayo, and O. Torres-Bugarín, "An update on peripheral ossifying fibroma: case report and literature review," *Oral and Maxillofacial Surgery*, vol. 20, no. 1, pp. 1–7, 2016.

[13] T. Farquhar, J. Macellan, H. Dyment, and Anderson R. D., "Peripheral ossifying fibroma: a case report," *Canadian Dental Association*, vol. 47, no. 9, 809 pages, 2016, 812.

[14] M. B. Mishra, K. A. Bhishen, and S. Mishra, "Peripheral ossifying fibroma," *Journal of Oral and Maxillofacial Pathology*, vol. 15, no. 1, pp. 65–68, 2011.

[15] A. Nekouei, A. Eshghi, P. Jafarnejadi, and Z. Enshaei, "A review and report of peripheral giant cell granuloma in a 4-year-old child," *Case Reports in Dentistry*, vol. 2016, Article ID 7536304, 4 pages, 2016.

[16] C. M. Flaitz, "Peripheral giant cell granuloma: a potentially aggressive lesion in children;", *Pediatric Dentistry*, vol. 22, no. 3, pp. 232–233, 2000.

[17] G. S. Letterman, "Peripheral giant cell granuloma," *Plastic and Reconstructive Surgery*, vol. 46, no. 3, p. 320, 1970.

[18] K. Veena, H. Jagadishchandra, J. Sequria, S. Hameed, L. Chatra, and P. Shenai, "An extensive denture-induced hyperplasia of maxilla," *Annals of Medical and Health Sciences Research*, vol. 3, no. 5, p. 7, 2013.

[19] E. M. Canger, P. Celenk, and S. Kayipmaz, "Denture-related hyperplasia: A clinical study of a Turkish population group;", *Brazilian Dental Journal*, vol. 20, no. 3, pp. 243–248, 2009.

[20] H. Mortazavi, H. R. Khalighi, S. Jafari, and M. Baharvand, "Epulis fissuratum in the soft palate: Report of a case in a very rare location," *Dental Hypotheses*, vol. 7, no. 2, pp. 67–69, 2016.

[21] I. M. Brook and D. J. Lamb, "Surgical/prosthetic problems of a large leaf fibroma," *Dental Update*, vol. 15, no. 3, p. 126, 1988.

[22] N. G. Nikitakis and J. K. Brooks, "Sessile nodule on the palate. Leaflike denture fibroma," *General Dentistry*, vol. 59, no. 1, pp. 76–77, 2011.

[23] B. Manovijay, P. Rajathi, S. M. Fenn, and B. Sekar, "Recurrent epulis granulomatosa: a second look," *Journal of Advanced General Dentistry*, vol. 59, no. 1, pp. 76–77, 2011.

[24] B. Manovijay, P. Rajathi, S. M. Fenn, and B. Sekar, "Recurrent epulis granulomatosa: a second look," *Journal of Advanced General Dentistry*, vol. 59, no. 1, pp. 76–77, 2011.
[25] S. Palanivelu, P. Jayanthi, U. K. Rao, E. Joshua, and K. Ranganathan, "Rapidly enlarging mass following dental extraction," Journal of Oral and Maxillofacial Pathology, vol. 15, no. 2, pp. 223–227, 2011.

[26] J. Faryabi and S. Adhami, "Unusual presentation of chronic hyperplastic pulpitis: a case report. Iran Endod J," Winter, vol. 2, no. 4, p. 156, 2008.

[27] N. S. A. Jabbar, J. M. Aldrigui, M. M. Braga, and M. T. Wanderley, "Pulp polypl in traumatized primary teeth: a case-control study," Dental Traumatology, vol. 29, no. 5, pp. 360–364, 2013.

[28] K. Anilkumar, S. Lingeswaran, G. Ari, R. Thyagarajan, and A. Logaranjani, "Management of chronic hyperplastic pulpitis in mandibular molars of middle aged adults: a multidisciplinary approach," Journal of Clinical and Diagnostic Research, vol. 10, no. 1, pp. ZD23–ZD25, 2016.

[29] M. Gorduysus, Hyperplastic Pulpitis Development in a Bridge Abutment Tooth, vol. 32, Hacettepe Dış Hekimliği Fakültesi Dergisi, Ankara, Turkey, 1 edition, 2008, 35–37.

[30] H. R. Khalighi, M. Hamian, F. M. Abbas, and S. Farhadi, "Simultaneous existence of giant cell fibroma and squamous papilloma in the oral cavity," Indian Journal of Medical Specialities, vol. 2, no. 2, 2011.

[31] N. G. Nikitakis, D. Emmanouil, M. P. Maroulakos, and M. V. Angelopoulou, "Giant cell fibroma in children: report of two cases and literature review," Journal of Oral and Maxillofacial Research, vol. 4, no. 1, article e5, pp. 1–7, 2013.

[32] V. K. K. Reddy, N. Kumar, P. Battapeti, L. Samyuktha, and S. P. Nanga, "Giant cell fibroma in a paediatric patient: a rare case report," Case Reports in Dentistry, vol. 2015, Article ID 240374, 3 pages, 2015.

[33] M. S. Thwaites, T. E. Jeter, and O. Ajagbe, "Inflammatory papillary hyperplasia: review of literature and case report involving a 10-year-old child," Quintessence International, vol. 21, no. 2, pp. 133–138, 1990.

[34] J. R. Antonelli, F. V. Panno, and A. Witko, "Inflammatory papillary hyperplasia: supraperiosteal excision by the blade-locating technique," General dentistry, vol. 46, no. 4, pp. 390–397, 1998.

[35] P. Infante-Cossio, R. Martinez-de-Fuentes, E. Torres-Carranza, and J. L. Gutierrez-Perez, "Inflammatory papillary hyperplasia of the palate: treatment with carbon dioxide laser, followed by restoration with an implant-supported prosthesis," British Journal of Oral and Maxillofacial Surgery, vol. 45, no. 8, pp. 658–660, 2007.

[36] H. Mohan, A. Tablan, I. Mundi, R. P. S. Punia, and A. Dass, "Non-neoplastic salivary gland lesions: a 15-year study," European Archives of Oto-Rhino-Laryngology, vol. 268, no. 8, pp. 1187–1190, 2011.

[37] H. Mortazavi, M. Baharvand, S. Alirezaei, and R. Noor-Mohammadi, "Combination therapy in a large lower lip mucoccele: a non-invasive recommended technique," Dental Hypotheses, vol. 5, no. 3, pp. 127–129, 2014.

[38] H. Mortazavi, H. R. Khalighi, M. Baharvand, M. Eshghpour, and R. Singh, "Bilateral symmetrical mucoccele of the lower lip: report of a rare clinical presentation," International Journal of Experimental Dental Science, vol. 3, pp. 92–94, 2014.

[39] R. Prakash, B. B. Kushwaha, and S. Gautam, "Airway management in a child with oral ranula," International Journal of Biomedical Research, vol. 5, no. 2, p. 148, 2014.

[40] S. Moghe, A. K. Pillai, S. Prabhu, S. Nahar, and U. K. Kartika, "Pleomorphic adenoma of the palate: report of a case," International Journal of Scientific Study, vol. 2, pp. 54–56, 2014.

[41] H. Mortazavi, S. Alirezaei, S. Azari-Marhabi, M. Baharvand, and M. Eshghpour, "Upper lip pleomorphic adenoma: comparison of reported cases between," Journal of Dental materials and Techniques, vol. 2, no. 4, pp. 125–129, 1990.

[42] M. Rahnama, U. Orzedala-Koszel, L. Czupkallo, and M. Lobacz, "Pleomorphic adenoma of the palate: a case report and review of the literature," Wspolczesna Onkologia, vol. 17, no. 1, pp. 103–106, 2013.

[43] S. I. Jarde, S. Das, S. Narayanswamy, A. Chatterjee, and C. Babu, "Mucoepidermoid carcinoma of the palate: a rare case report," Journal of Indian Society of Periodontology, vol. 20, no. 2, pp. 203–206, 2016.

[44] P. Ritwik, K. G. Cordell, and R. B. Brannon, "Minor salivary gland mucoepidermoid carcinoma in children and adolescents: a case series and review of the literature," Journal of Medical Case Reports, vol. 6, article no. 182, 2012.

[45] N. Shah, A. Mahjan, H. Patel, R. Shah, and S. Shah, "Mucoepidermoid carcinoma of palate: a case report," Scholars journal of dental sciences, vol. 2, no. 3, pp. 222–224, 2015.

[46] K. Triantafillidou, J. Dimitrakopoulos, F. Iordanidis, and D. Koufogiannis, "Management of adenoid cystic carcinoma of minor salivary glands," Journal of Oral and Maxillofacial Surgery, vol. 64, no. 7, pp. 1114–1120, 2006.

[47] P. J. Bradley, "Adenoid cystic carcinoma of the head and neck: a review," Current Opinion in Otolaryngology & Head and Neck Surgery, vol. 12, no. 2, pp. 127–132, 2004.

[48] M. J. Mendez, V. C. Carrard, A. N. Haas et al., "A 10-year study of specimens submitted to oral pathology laboratory analysis: lesion occurrence and demographic features," Brazilian Oral Research, vol. 26, no. 3, pp. 235–241, 2012.

[49] M. Ali and D. Sundaram, "Biopsied oral soft tissue lesions in Kuwait: a six-year retrospective analysis," Medical Principles and Practice, vol. 21, no. 6, pp. 569–575, 2012.

[50] P. Dalvi, K. Vandana, S. Prakash, and K. Mohan, "Solitary neurofibroma of the gingiva: a rare case report," Indian Journal of Multidisciplinary Dentistry, vol. 6, no. 2, p. III, 2016.

[51] J. P. George and N. Sai Jyothsna, "Solitary neurofibroma: a rare occurrence on gingiva," General Dentistry, vol. 64, no. 3, pp. 28–31, 2016.

[52] T. Shimoyama, T. Kato, D. Nasu, T. Nakano, N. Horie, and F. Ide, "Solitary neurofibroma of the oral mucosa: a previously undescribed variant of neurofibroma," Journal of oral science, vol. 44, no. 1, pp. 59–63, 2002.

[53] P. Prasanna Kumar and K. Meghashri, "Schwannoma of the hard palate: a case report and review of literature," Journal of Oral Research and Review, vol. 3, no. 1, pp. 23–27, 2012.

[54] J.-M. Sanchis, C.-M. Navarro, J.-V. Bagà et al., "Intraoral schwannomas: presentation of a series of 12 cases," Journal of Clinical and Experimental Dentistry, vol. 5, no. 4, pp. 192–196, 2013.

[55] S. Parhar, H. Preet Singh, A. Nayyar, and A. S. Manchanda, "Intra-oral schwannoma: a case report," Journal of Clinical and Diagnostic Research, vol. 8, no. 3, pp. 264–265, 2014.

[56] S. Kumaraswamy, N. Madan, R. Keerthi, and S. Shakti, "Lipomas of oral cavity: case reports with review of literature," Journal of Maxillofacial and Oral Surgery, vol. 8, no. 4, pp. 394–397, 2009.
[57] R. Kaur, S. Kler, and A. Bhullar, “Intraoral lipoma: report of 3 cases,” Dental Research Journal, vol. 8, no. 1, pp. 48–51, 2011, (Isfahan) Winter.

[58] K. Rana, V. Narula, E. K. Bhargava, R. Shankar, and N. Mahajan, “T-cell lymphoma of the oral cavity: case report,” Journal of Clinical and Diagnostic Research, vol. 9, no. 3, pp. MD03–MD04, 2015.

[59] M. Malaguarnera, M. Giordano, C. Russo et al., “Lymphoma with sclerotherapy: case report,” Journal of Oral Pathology and Medicine, vol. 40, no. 2, pp. 167–173, 2011.

[60] M. Kaur, N. Kler, and A. Bullar,”Introral lipoma: report of two cases,” Journal of Oral Pathology and Medicine, vol. 3, no. 1, pp. 116, 2012.

[61] L. A. Goodstein, A. Khan, J. Pinczewski, and V. N. Young, “Orallymphangioma:ararecasereport,” Jornal Vascular Brasileiro, vol. 13, no. 3, 2014.

[62] A. Dilsiz, T. Aydin, and N. Gursan, “Capillary hemangioma as a rare benign tumor of the oral cavity: a case report,” Cases Journal, vol. 2, no. 9, article no. 8622, 2009.

[63] K. A. Kamala, L. Ashok, and G. P. Sujatha, “Cavernous hemangioma of the tongue: a rare case report,” Contemporary Clinical Dentistry, vol. 5, no. 1, pp. 95–98, 2014.

[64] V. Usaha, T. Sivasankari, S. Jeeani, G. S. Asokan, and J. Parthiban, “Lymphangioma of the tongue: a case report and review of literature,” Journal of Clinical and Diagnostic Research, vol. 8, no. 9, pp. ZD12–ZD14, 2014.

[65] H. Bhayya, D. Pavan, M. L. Avinash Tejasvi, and P. Geetha, “Oral lymphangioma: a rare case report,” Contemporary Clinical Dentistry, vol. 6, no. 4, pp. 584–587, 2015.

[66] S. Sunil, D. Gopakumar, and B. Sreenivasan, “Oral lymphangioma: case reports and review of literature,” Contemporary Clinical Dentistry, vol. 3, no. 1, p. 116, 2012.

[67] H. Alan, S. Agacayak, G. Kavak, and A. Ozcan, “Verrucous carcinoma and squamous cell papilloma of the oral cavity: report of two cases and review of literature,” European Journal of Dentistry, vol. 9, no. 3, pp. 453–456, 2015.

[68] L. A. Goodstein, A. Khan, J. Pinczewski, and V. N. Young, “Symptomatic squamous papilloma of the uvula: report of a case and review of the literature,” Case Reports in Otolaryngology, vol. 2012, pp. 1-2, 2012.

[69] P. P. Jaju, P. V. Suvarna, and R. S. Desai, “Squamous papilloma: case report and review of literature,” International Journal of Oral Science, vol. 2, no. 4, pp. 222–225, 2010.

[70] M. Nagaraj, “Verruca vulgaris of the tongue,” Journal of Maxillofacial and Oral Surgery, vol. 12, no. 3, pp. 329–332, 2013.

[71] A. Ural, S. Arslan, Ş. Ersoz, and B. Değer, “Verruca vulgaris of the tongue: a case report with literature review,” Bosnian Journal of Basic Medical Sciences, vol. 14, no. 3, pp. 136–138, 2014.

[72] H. Mortazavi, M. Baharvand, and M. MehdiPour, “Oral potentially malignant disorders: an overview of more than 20 entities,” Journal of Dental Research, Dental Clinics, Dental Prospects, vol. 8, no. 1, p. 14, 2014, Winter.

[73] M. Zargaran, N. Eshghyar, P. B. Vaziri, and H. Mortazavi, “Immunohistochemical evaluation of type IV collagen and laminin-332 γ2 chain expression in well-differentiated oral squamous cell carcinoma and oral verrucous carcinoma: a new recommended cut-off,” Journal of Oral Pathology and Medicine, vol. 40, no. 2, pp. 167–173, 2011.