Proliferative Verrucous Leukoplakia - A Clinico Pathological Entity: An update

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ABSTRACT:

Proliferative verrucous leukoplakia (PVL) is considered to be a progressive type of oral leukoplakia that is impossible to recognize in its nascent stages. The condition typically begins as a white plaque of hyperkeratosis that eventually becomes a multifocal disease with confluent exophytic and proliferative features. It is observed most frequently in elderly women of fifth and six decades of life at the time of diagnosis. This article describes the clinical aspects and histological features and dealing with controversies in diagnosis.

Key words: leukoplakia, exophytic, proliferative

the vestibule), tongue, and buccal mucosa. PVL often begins as a focal lesion spreading laterally over time and can be multifocal. Early in its course it is a flat hyperkeratotic lesion that becomes progressively verrucous and histologically often exhibits varying degrees of epithelial dysplasia.4

Case Report:

A 59-year-old female patient reported to the Department of Oral Medicine and Radiology with the chief complaint of rough white patch on right alveolar mucosa since 18 months. She gave a history of chutta smoking since 40yrs. Intra oral examination revealed white verrucous slightly raised lesion with granular texture measuring 4 x 2cms on right alveolar mucosa (Figure 1). The lesion was extending anterio posteriorly from mesial aspect of 47 to anterior mandibular right alveolar ridge. Bucco-lingually involves entire mandibular right alveolar ridge and extends to vestibule upto mesial aspect of 47. On palpation the growth was firm, non tender, non fluctuant and non compressible. Extra oral examination revealed right submandibular lymphadenopathy which was non tender and mobile. Based on the history and clinical examination a Provisional diagnosis of Proliferative verrucous leukoplakia was made with a differential diagnosis of Candidal leukoplakia, and verrucous carcinoma. The patient was subjected to following investigations to reach to a final diagnosis Orthopantomograph (OPG), Complete hemogram, Incisional biopsy. Orthopantomograph reveals severe bone loss in relation to 47 and also showing that all remaining right mandibular teeth were missing. In complete hemogram all the values were in the normal range. An incisional biopsy was performed from right alveolar mucosa and the tissue was sent for histopathological analysis of mandibular multiple missing teeth and an incisional biopsy of the lesion was done for confirmation.

Histopathology showed marked hyperkeratosis with pseudoepitheliomatous hyperplasia with focal connective tissue entrapment. The surface of epithelium shows exophytic type of growth pattern with Parakeratin plugging. There is evidence of keratin pearl formation in few areas of the epithelium. Sub epithelial connective tissue shows mild lymphocytic infiltrate with no evidence of malignancy and was diagnosed as verrucous hyperplasia (Figure 2). Correlating clinical and histological features final diagnosis of Proliferative verrucous leukoplakia was given. The entire lesion was surgically excised. Later the follow-up was not possible as patient did not report back.

Discussion:

According to the latest WHO nomenclature, proliferative verrucous leukoplakia conforms to the new terminology of “potentially malignant disorders” given that it is neither a Delimited lesion nor a condition.1 PVL presents with specific characteristics, mainly a more aggressive biological behavior than other forms of leukoplakia expressed by: A tendency toward multifocality; a high probability of recurrence; and a high rate of malignant transformation, which can range between 40% and 100% in a follow up period of 4.4 to 11.6 years.3 The lesions are slow growing yet persistent, as well as irreversible and resistant to all forms of treatment with a high recurrence rate. Throughout its development, it is common to find erythematous and/or verrucous areas that occasionally progress to verrucous carcinoma or squamous cell carcinoma. PVL is more common in elderly women who have had lesions of leukoplakia for many years. The mean age at the time of diagnosis is over 60 years.6 In the present case the patient was 59 yrs female and the growth was verrucous.

Etiopathogenesis:

PVL is of uncertain etiology. An association with human papilloma virus(HPV) infection, particularly 16 and 18 has been suggested.7 Palefsky et al. (1995) studied nine lesions from seven patients with PVL and they found that eight lesions(89%) were positive for HPV and out of these eight, seven contained HPV 16. They therefore suggested that HPV 16 infection may play an important role in occurrence of these lesions.8 Aberrations in the cell cycle regulatory genes p16INK4a and p14ARF with homozygous deletion, loss of heterozygosity, and mutation appear to be frequent findings in PVL.8

Histopathological features:

The histologic findings are variable and usually a function of time or age of the lesion. The early lesions are deceptively bland exhibiting only hyperkeratosis, but over time they become progressively verrucous and often show varying degrees of epithelial dysplasia. The advanced lesions have a propensity for conventional squamous cell carcinoma and verrucous carcinoma within the histologic spectrum.9
Hansen et al suggested histological stages in the continuum of PVL with intermediates as follows:

**Grade 0**: Normal mucosa; **Grade 2**: Hyperkeratosis (clinical leukoplakia); **Grade 4**: Verrucous hyperplasia; **Grade 6**: Verrucous carcinoma; **Grade 8**: Papillary squamous cell carcinoma; **Grade 10**: Less well differentiated squamous cell carcinoma.

Batsakis et al reduced the number of histological stages to four with intermediates.  
**Grade 0**: Clinical flat leukoplakia without dysplasia; **Grade 2**: Verrucous hyperplasia; **Grade 4**: Verrucous carcinoma; **Grade 6**: Conventional squamous cell carcinoma with intermediates.

**Diagnosis:**

Because of the lack of specific histological criteria, the diagnosis of PVL is based on combined clinical and histopathologic evidence of progression. In previously published series, diagnosis of PVL was made according to Hansen’s definition. There are few studies that apply a set of diagnostic criteria that are mentioned as follows:

Ghazali et al., established the following criteria:

- The lesion starts as homogenous leukoplakia without evidence of dysplasia at the first visit
- With time, some areas of leukoplakia become verrucous
- The disease progresses to the development of multiple isolated or confluent lesions at the same or a different site with time
- The disease progresses through the different histopathological stages reported by Hansen et al
- The appearance of new lesions after treatment
- A follow up period of no less than one year
- Diagnosis of PVL changes gradually from a simple plaque of hyperkeratosis without dysplasia to verrucous hyperplasia, verrucous carcinoma or OSCC.

Cerero Lapiedra et al established the following major and minor criteria:

**Major criteria:**

(a) Leukoplakia lesion with more than two different oral sites, which is most frequently found in the gingiva, alveolar processes and palate
(b) The existence of a verrucous area
(c) That the lesions have spread or engrossed during development of the disease
(d) That there has been a recurrence in a previously treated area
(e) Histopathologically, there can be from simple epithelial hyperkeratosis to verrucous hyperplasia, Verrucous carcinoma or OSCC, whether in situ or infiltrating.
Minor criteria:
(a) An OL lesion that occupies at least 3 cm when adding all the affected areas
(b) That the patient be female
(c) That patient (male or female) be a non smoker
(d) A disease evolution higher than 5 years.

In order to make the diagnosis of PVL, it was suggested that one of the two following combinations of the criteria mentioned before were met. Three major criteria (E being among them) or Two major criteria (E being among them) + two minor criteria. Nevertheless, at present, there is no criterion that will allow for the early diagnosis of the disease.

In present case it was fulfilling 4 major criteria and 2 minor criteria.

Treatment:
The lesions of PVL are persistent, progressive, and relentless and have a high recurrence rate regardless of the treatment method employed. Complete surgical excision is difficult because of the diffuse, multifocal nature of the condition and the inability to assess accurately surgical margins clinically and microscopically. Other modalities, such as laser ablation, external beam radiation, and chemotherapy, have been unsatisfactory. Methisoprinol as a synthetic agent capable of inhibiting viral ribonucleic acid synthesis and replication and of stimulating antiviral cell mediated reactions that have been shown to have some clinical efficacy in HPV - induced lesions. Topical photodynamic therapy also may prove useful. It causes relatively low morbidity ad no scarring and multiple mucosal sites can be treated simultaneously.13

Malignant transformation and recurrences:
PVL is characterized not only by a high rate of recurrences after treatment but also by malignant transformation in nearly 74% of cases, with a tendency for several oral cancers to appear. The gingiva and palate represented the areas with the highest frequency of these multiple malignant tumor.14

Conclusion:
The biologic behavior of PVL is decidedly more aggressive than the other forms of leukoplakia. However, the initial clinical presentation of PVL is so innocuous that one can easily dismiss it as insignificant until there is widespread oral involvement and carcinomatous transformation. Therefore, it is recommended to have the earliest possible diagnosis, as well as consensus on diagnostic criteria to achieve uniformity. To this effect, the present article mentioned a set of diagnostic criteria, which we believe should be applied to case series in order to verify its efficacy.

References: