Premalignant and Malignant Lesions of Oral Cavity

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Anatomy

- The palate forms the roof of the mouth and intervenes between the nasal and oral cavities.
- It consists of the palatine process of the maxilla, the horizontal plates of the palatine bone.
Three foramina open on the oral aspect of the hard palate

- **Incisive Fossa**
  - Slight depression posterior to central incisor teeth
  - Nasopalatine nerve
- **Greater palatine foramina**
  - Medial to 3rd Molar
  - Greater palatine vessels and nerve
- **Lesser palatine foramina**
  - Lesser Palatine nerves and vessels to soft palate
Blood Supply

- Greater palatine artery
- Superior Alveolar Arteries
- The Greater Palatine artery is a branch of the third part of the maxillary artery. The greater palatine artery descends with its accompanying nerve in the palatine canal.
- The superior alveolar arteries are terminal branches of the nasopalatine artery
The greater palatine emerges on the hard palate from the greater palatine foramen runs forward in a groove on the inferior surface of the bony palate almost to the incisor teeth supplies the gums and the mucosa and glands of the hard palate.
The venous drainage is to the pterygoid plexus and subsequently to the internal jugular venous system.
Innervation

- The nasopalatine nerves
- Greater Palatine Nerves
- The nasopalatine nerves are branches of the maxillary division of the trigeminal nerve.
- They enter the palate at the incisive foramen supply the anterior part of the hard palate behind the incisor teeth.\(^5\)
• Greater (and Lesser) Palatine run through the palatine canal and exit at the Great and Lesser Palatine Foramens, respectively.

• Parasympathetic postganglionic secretomotor fibres from the pterygopalatine ganglion run with the nerves to supply the palatine mucous glands.
Perineural Spread

- Tumors spreading by perineural extension can be discovered by radiographic enlargement of the palatine foramina or widening of the palatine canals or the foramen rotundum.\textsuperscript{6}
Oral submucous fibrosis (OSMF)

- OSMF is a high risk precancerous condition that predominantly occurs amongst Indians.
- Factors implicated in the pathogenesis of sub mucous fibrosis:
  - Chillie consumption
  - Areca-nut chewing, autoimmunity
  - Genetic predisposition
- Now there is convincing epidemiologic evidence implicating areca nut as a causative factor in its pathogenesis.
Epidemiology

- The prevalence of OSMF in random samples of the population in India is up to 0.4%.
- Although hard data are not available, indications are that this disease is increasing rapidly in India.
- Submucous fibrosis occurs in both sexes over a wide age range.
Definition

- OSMF is a chronic mucosal condition affecting any part of the oral mucosa.
- **Mucosal rigidity** of varying intensity due to fibroelastic transformation of the juxtaepithelial connective tissue layer.
- The presence of palpable fibrous bands is a diagnostic criterion for submucous fibrosis.
• When the **tongue** is affected, it is devoid of papillae and its mobility, especially the protrusion, is **impaired**
• The **opening of the mouth** is restricted
• In severe OSMF, the patient cannot protrude the tongue beyond the incisal edges and there is a progressive closure of the oral opening
• OSMF must be diagnosed only if palpable fibrous bands are present
Clinical aspects

- The most common initial symptoms:
  - Burning sensation of the oral mucosa aggravated by spicy food followed by either hypersalivation or dryness of the mouth.
• The most common and initial clinical sign as well as a regular feature:
  ▫ blanching i.e., marble-like appearance of the oral mucosa.
• In advanced cases, the mucosa becomes tough and leathery, with numerous vertical fibrous bands
Natural History

• Unlike precancerous lesions OSMF is not known to regress, either spontaneously, or with the cessation of the areca-nut chewing habit.

• The most serious aspect of this disease is the high risk for the development of oral cancer.

• The epithelium is atrophic in this condition which renders it susceptible to the action of carcinogens.
• OSMF and coexistent leukoplakia: Leukoplakia is a precancerous lesion; its coexistence with OSMF implies the high risk for oral cancer.

• OSMF and coexistent oral cancer: Not uncommonly (in 5% to 42% of the cases), submucous fibrosis and oral cancer coexist

• Malignant transformation: Long-term population based studies have confirmed its precancerous nature.
• No definitive and widely accepted treatment is currently available.
• Some temporary relief from the symptoms and improvement in the oral opening with medicinal treatment such as local injections of cortisone and placentrex.
• It is essential to follow-up the patients regularly.
• Patient education to discontinue the use of areca nut and tobacco in any form.
LEUKOPLAKIA
• Leukoplakia is the most common premalignant or "potentially malignant" lesion of the oral mucosa
• It is a predominantly white lesion of the oral mucosa
• The incidence and prevalence of leukoplakia vary in different parts of the world
In general the reported prevalence ranges from 0.2 to 5% (India 0.2-4.9%). It is seen most frequently in middle-aged and older men. Gender distribution is also variable. Men are more affected in some countries.
Clinical presentation

- Leukoplakia can be either solitary or multiple
- It may appear on any site of the oral cavity, the most common sites being: buccal mucosa, alveolar mucosa, floor of the mouth, tongue, lips and palate
Classically two clinical types of leukoplakia are recognised

- **Homogeneous leukoplakia** is defined as a predominantly white lesion of uniform flat and thin appearance that may exhibit shallow cracks. This type is usually asymptomatic.

- **Non-homogeneous leukoplakia** has been defined as a predominant white or white-and-red lesion ("eritroleukoplakia") that may be either irregularly flat, nodular ("speckled leukoplakia") or exophytic ("exophytic or verrucous")
Homogeneous leukoplakia on the dorsum and left lateral margin of the tongue
Non-homogenous Leukoplakia
Verrucous Leukplakia
Aetiopathogenesis

- The aetiology of leukoplakia is still unclear
- Tobacco seems to be the major inductor factor, its association cannot be determined in all cases
- A variety of smokeless tobacco habits have been reported as leukoplakia inductors: e.g. snuff, chewing. These lesions have shown to have a low malignant transformation risk
Other factors such as:

- Alcohol
- Inadequate diet
- Vitamin deficiency (e.g. vitamin A and C), areca nut (betel)
- Chronic traumatic irritation
- Poor oral hygiene
- Poor socio-economic status.
Treatment

• There are different treatments for leukoplakia.
• However, the risk of malignant transformation is not completely eliminated by any of the current therapies.
• Initial treatment of a white oral lesion is the elimination of the possible aetiological factors.
• Complete surgical removal (leaving free-lesion borders) is recommended in cases with epithelial dysplasia.
• Apart from surgical excision, other treatment modalities available include cryosurgery, laser surgery, retinoids, beta-carotene, bleomycin, calcipotriol, photodynamic therapy.
Prognosis and complications

- The malignant transformation rate of oral leukoplakia varies from 0 to 33%.
- Regular check-up of these patients is essential, probably every 3, 6 and then 12 months, both in treated and untreated patients.
Oral Cancer
Oral Cancer is the sixth leading cause of cancer worldwide.
The survival rate was 52%.
Oral cancer generally are socially derived diseases.
Tobacco and alcohol have synergistic effect.
Treatment of early oral cancer is surgery. Locally advanced T3/4 are best treated with combined surgery and Radiotherapy.
High risk of second primary cancer.
EPIDEMIOLOGY

- The Oral cavity extends from vermilion border of lips to the plane between junction of the hard palate and soft palate.
- Include: Lips and oral cavity (buccal mucosa, tongue, gingiva, retromolar trigone, flour of mouth, hard palate)
- The incidence of oral cancer varies throughout the world.
  High incidence in India, France, SE Asia.
- 40% of HN cancer
- Age onset 50 yrs. Sex ratio 3:1
Risk factors

- Heavy tobacco
- Alcohol.
- Syphilis
- Viruses (EB, HSV, HPV, HIV)
- Neglect of oral dental hygiene (chronic infection, unfit dentures)
- Lichen planus, Plummer Vinson sy.
- Immunosuppression, malnutrition
Those who use tobacco and alcohol simultaneously are thought to have a significantly increased risk of oral cancer relative to using either one alone, likely because the combined use of nicotine and ethanol (known cytotoxins) significantly increases the penetration of N-Nitrosonornornicotine (NNN), a known carcinogen found in tobacco, across the oral mucosa.
Distribution of Oral Cancer According to Locations

- Tongue: 34%
- Retromolar: 2%
- Ginggiva Max: 12%
- Ginggiva Mand: 7%
- Buccal: 24%
- Lower Lip: 16%
- Palatum: 5%
- Ginggiva Max: 12%
Pathology

- 90% SCC: Well/Moderate/Poorly/Undiff
- Exophytic, Ulcerative, Infiltrative, verrucous
- Other: Adeno Ca / from malignant minor salivary gland tumors, Melanoma, Sarcomas.
- Premalignant lesions: Leucoplakia, hyperplasia, Erythroplakia, and dysplasia
- Regional Lnn meta related to size and thickness of primary tumor
Clinical presentation

• Non healing ulcers
• Induration
• Verucous/cauliflower
• Hot potato chewing
• Trismus
• Lnn enlargement
Soft Palate vs. Hard Palate

Tumors of the soft palate
• similar to other tumors of the oral cavity
  ▫ 90% of all oral cancer is squamous cell\textsuperscript{1}
• 69.7% of soft palate tumors are squamous cell carcinoma\textsuperscript{2}

Tumors of the hard palate
• Squamous:Non-Squamous is 1:2 to 1:4
• Varied histology with non-squamous cell tumors,
  ▫ minor salivary gland tumors
  ▫ rare cases of melanoma\textsuperscript{3}
  ▫ sarcoma
  ▫ malignant lymphomas\textsuperscript{4}
Diagnosis

- Clinical:
  - History
    Detail clinical examination (used head lamp, mirror)
    Bimanual palpation
  - Cervical Lnn examination
  - Endoscopy (searching the second primary)
  - Biopsy
  - Staging: Panoramic photo, thorax, USG liver, or CT/MRI/PET Scan
TREATMENT

• Treatment Goals:
  To eradicate the primary tumor and LN metastasis, to maintain the function, and cosmetic reconstruction.

• Factors affecting choice of treatment:
  Tumor factors
  Patient factors
  Resource factors
TREATMENT

• SURGERY:
  Early stage T1/2No tumor: Wide excision +/- ND
  High risk of locoregional recurrent (40%)

• Management of No Neck:
  High incidence of occult metastasis in the clinically No Neck (15-43%)
  Controversy: Observation or Surgery/Radiation
  Depend on primary site.
  Should have minimal morbidity
  ELND if risk of occult meta >20%. (SND/SOHND).
  Sentinel Lymph Node Biopsy (SLNB)?

• Locally advanced tumor: Combined modality treatment
Tumour factors
- Site
- Size
- Location (anterior v/s posterior)
- Proximity to bone
- Status of cervical lymph nodes
- Previous treatment
- Tumour grade
- Depth of invasion

Patient factors
- Age
- General medical condition
- Performance status
- Occupation
- Lifestyle (smoking/drinking)
- Socio-economic considerations
- Previous treatment

Physician factors
- Surgery
- Radiotherapy
- Chemotherapy
- Nursing & rehabilitation services
- Dental
- Prosthetics
- Support services
6 Levels of Lymph-Nodes
Selective Neck Dissection
Classification of ND

1991 Classification:
- RND
- Modified RND
- Selective ND:
  - Supraomohyoid
  - Lateral
  - Posterolateral
  - Anterior
- Extended ND

2001 Classification:
- RND
- Modified RND
- Selective ND (SND):
  - SND (L.I-III/IV)
  - SND (L.II-IV)
  - SND (L.II-V)
  - SND (L.VI)
- Extended ND

Proposed by American HN Society and AAOHNS
Selective Neck Dissection
Modified RND 123
SURGICAL APPROACHES

- Trans-oral approach
- Lower cheek approach
- Upper cheek approach
- Swing mandibulotomy
- Visor flap
RECONSTRUCTION

• Single-stage immediate reconstruction is recommended.
• The technique:
  Skin grafts
  Pedicle flaps
  Alloplastic materials
  Autografts
  Free flaps
Adjuvant treatment

- Radiotherapy (External beam/Interstitial)
- Chemotherapy
- Concomittant Radio+Chemotherapy (Neoadjuvant)

- Palliative Chemotherapy for advanced diseases
PROGNOSIS

- Location/thickness/depth of primary tumor
- Staging
- Type of histology
- Grading
- Presence of perineural spread
- Mandibular invasion
- Lnn extension (Level, size, extracaps of meta)
- Molecular markers (?)
Summary

• The main problem of oral cancer is early detection
• Surgery is still the most important modality in management of oral cancer.
• Better understanding of molecular biology of HNSCC.
• Bio-molecular markers can be used in the management of SCC oral cancer.
• High risk of second primary cancer, Chemoprevention?