

بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ

Cementum and Alveolar bone

• أعداد:

• د. نور صباح أرحيم

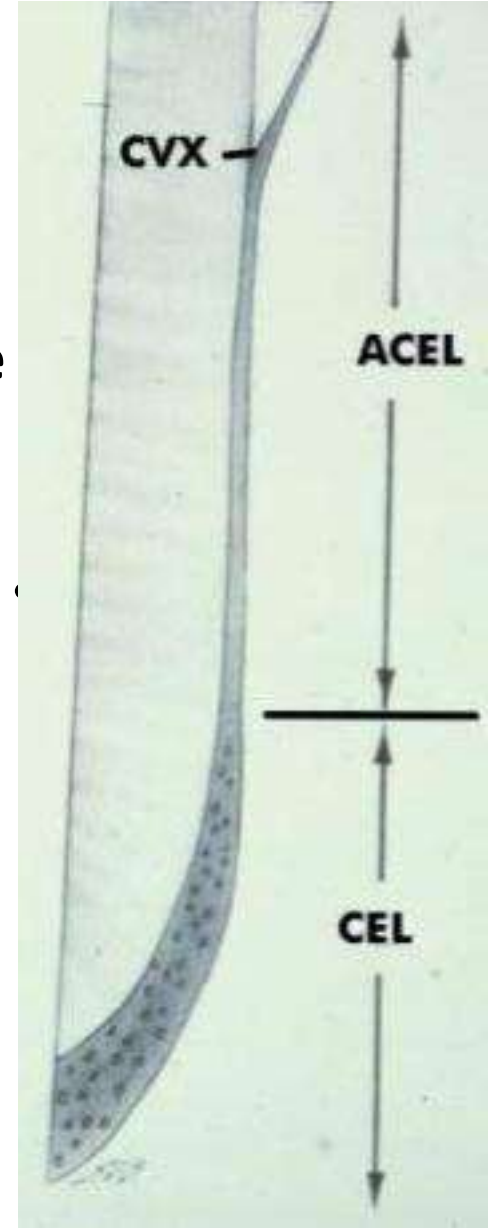
Cementum:

➔ A thin specialized calcified tissue covering the roots surfaces of the teeth.

•

➔ There are two main types of root cementum : acellular (primary) and cellular (secondary) .

•



Functions of cementum:

- 1- Anchorage of the tooth in the alveolus. •**
- 2-To attach the PDL fibers to the teeth. •**
- 3- To contribute to the process of repair after damage to the root surface and following regenerative periodontal surgical procedures. •**

Structures of cementum: cementum consist of :

1- Fibrous elements (collagen fibers).

- **Extrinsic fibers (Sharpey's fibers).**

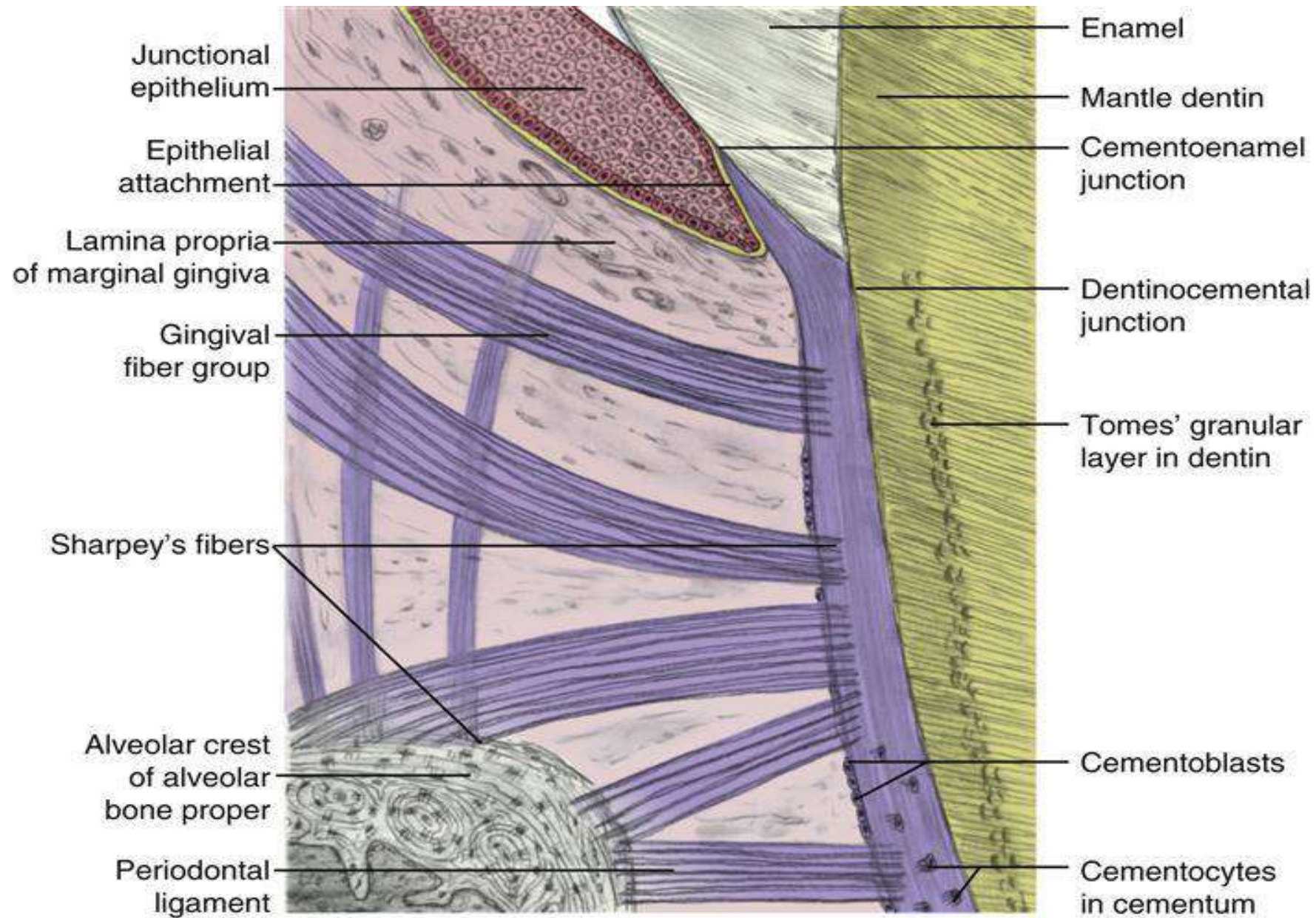
- **Intrinsic fibers.**

2- Cellular elements (cementoblast, cementocyte, fibroblast, cementoclast) .

3- Calcified interfibrillar matrix (proteoglycans, glycoprotein, phosphoprotein).

Extrinsic fibers: (Sharpey's fibers): which are the embedded portion of the principal fibers of the PDL and are formed by the fibroblast cells. Sharpey's fibers make up most of the structure of acellular cementum and they are inserted at right angles to the root surface and penetrate deep into the cementum.

Intrinsic fibers: these fibers are produced by cementoblast cells and are oriented more or less parallel to the long axis of the root and form a cross-banding arrangement with Sharpey's fibers.



1- cementoblast: responsible for the formation of both cellular and acellular cementum.

2-cementocyte: are found only in cellular cementum , they are located within spaces (lacunae) that communicate with each other through canaliculi for transportation of nutrition through the cementum and contribute to maintenances of the vitality of this tissue.

3-fibroblast: these are belong to the PDL where they are responsible for the synthesis of principle fibers but since these fibers become embedded in cementum, fibroblasts indirectly particitate in the formation of cementum.

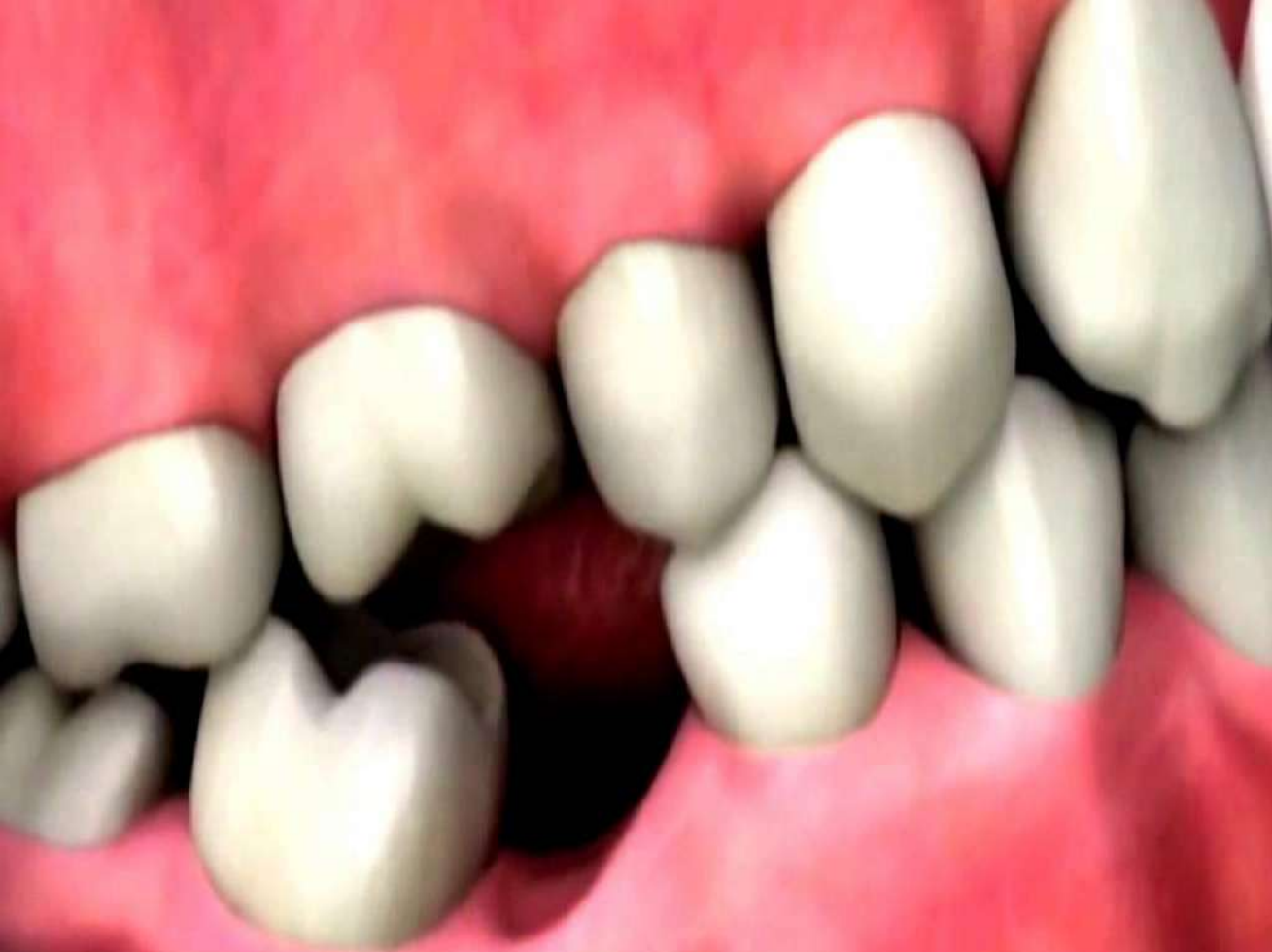
4-cementoclast: these cells are responsible for extensive root resorption that lead to primary teeth exofoliation . Permenant teeth do undergo physiological resorption but localized cemental resorption may occur which appear as concavities in the root surface and may be caused by local or systemic causes: **local condition** include, trauma from occlusion, orthodontic movement , cyst., while in **systemic condition** caused in calcium deficiency and hypothyroidisms.

Mineralization of cementum:

- ★ Occur by the deposition of hydroxyapatite crystals, first within the collagen fibers, later upon the fiber surface and finally in the interfibrillar matrix.
- ★ cellular cementum is less calcified than acellular cementum.
- ★ cementum mineralization is less than that of the bone, enamel and dentin.

Development of cementum:

- 1- Both cellular and acellular cementum are produced by cementoblast cell.
- 2-cementoid is first formed which is a non calcified tissue containing collagen fibrils distributed in matrix.
- 3-cementum thickness increase with age.
- 4- cementum thickness is more in apical area and furcation area than cervical area.
- 5-cementum is thicker in distal surface than in mesial surface because of functional stimulation from mesial drift.



And the opposing tooth begins to extrude



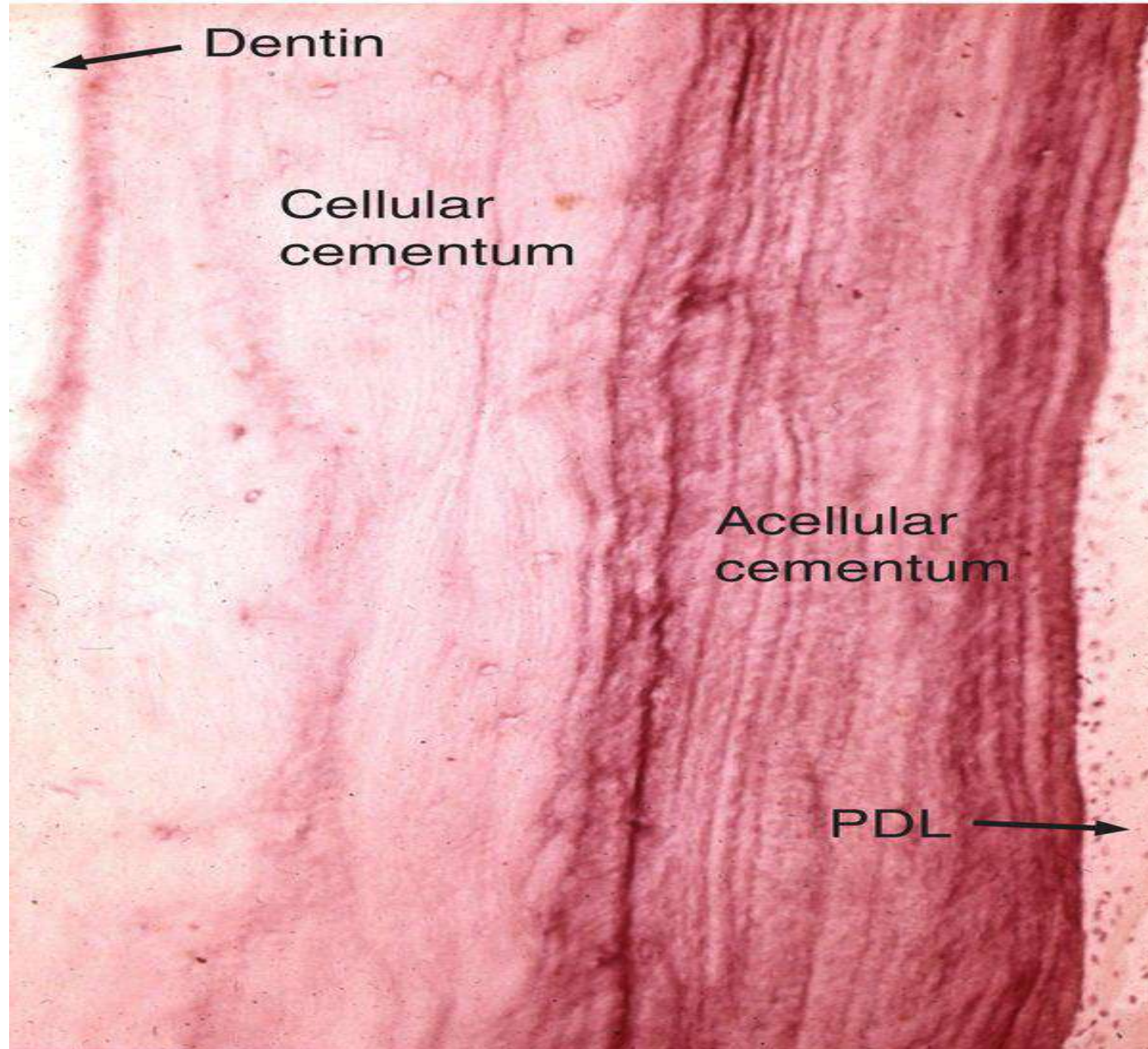
There are two types of cementum(both produced by cementoblast cells):

1- Primary (acellular cementum).

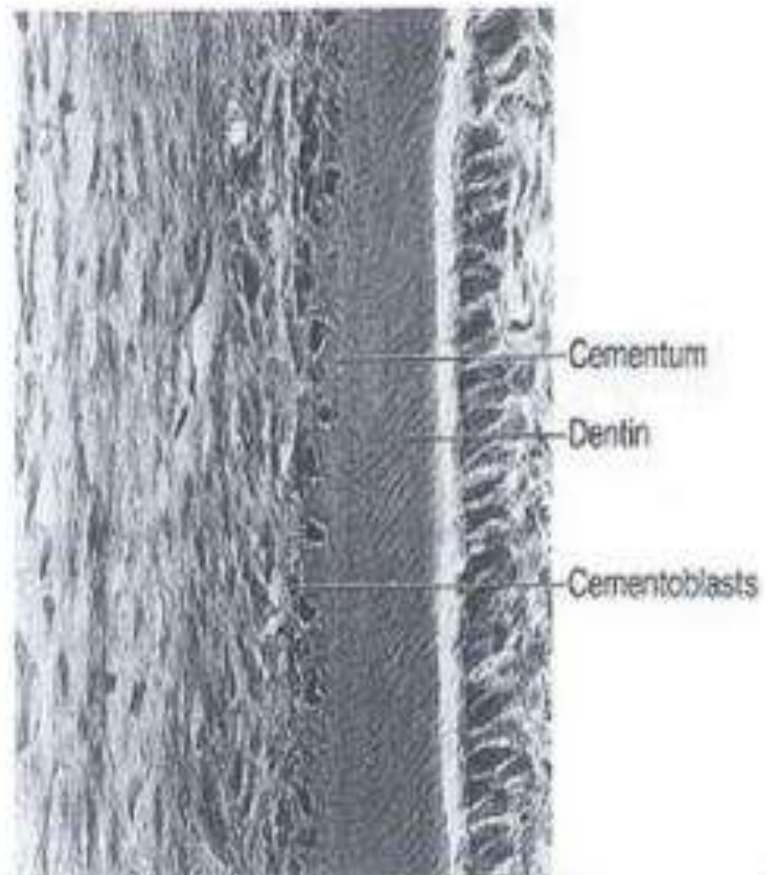
- First to be formed in conjunction with root formation and tooth eruption.**
- it is not contain cells and sharpys fibers make up most of its structures.**
- It cover the cervical third of the root.**

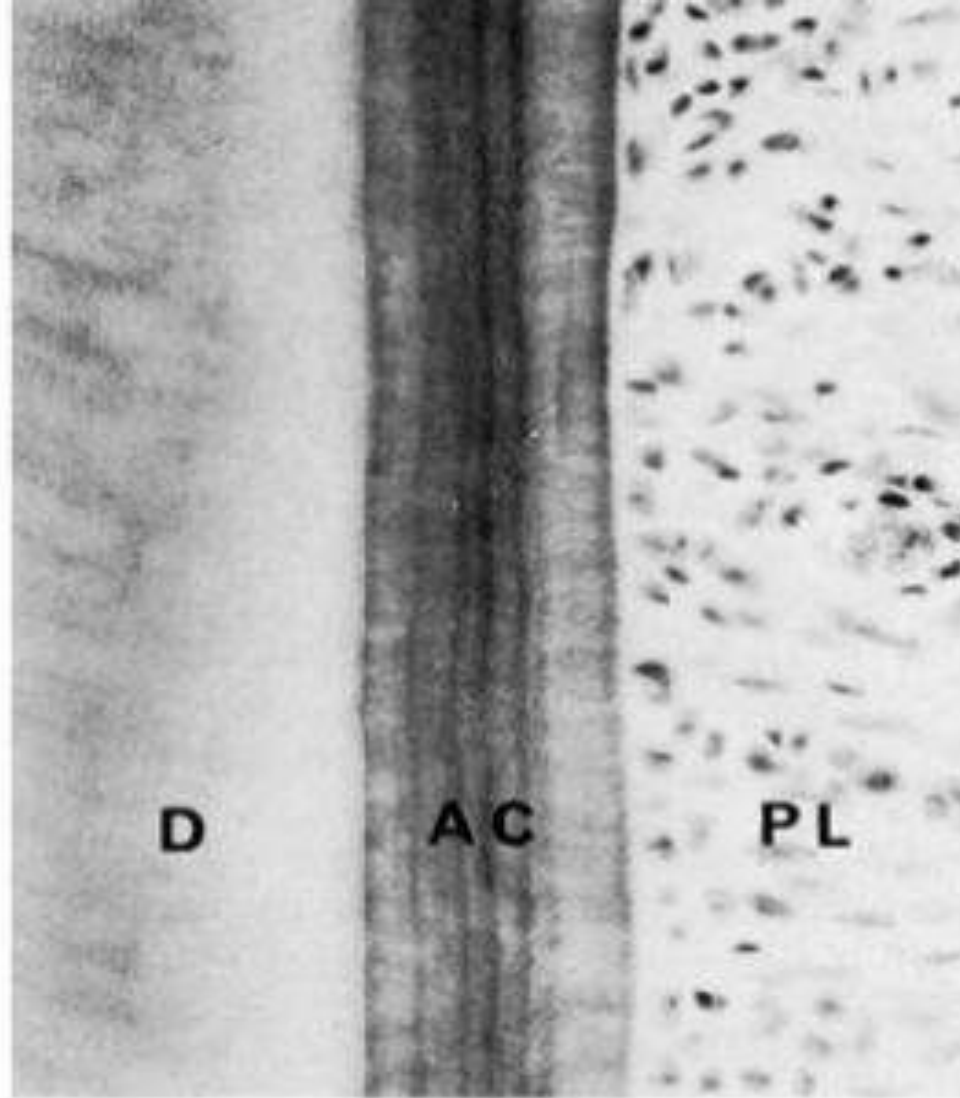
2- Secondary (cellular cementum).

- formed after tooth eruption in response to functional demand.**
- Grow faster and over a thin layer of acellular cementum at the apical third of the root and furcations of multirouted teeth.**
- Contain cells(cementocyte), Sharpys fibers occupy small portion.**
- Is less calcified than a cellular type.**



CELLULAR CEMENTUM





Copyright © 2006 by Saunders, an imprint of Elsevier Inc.

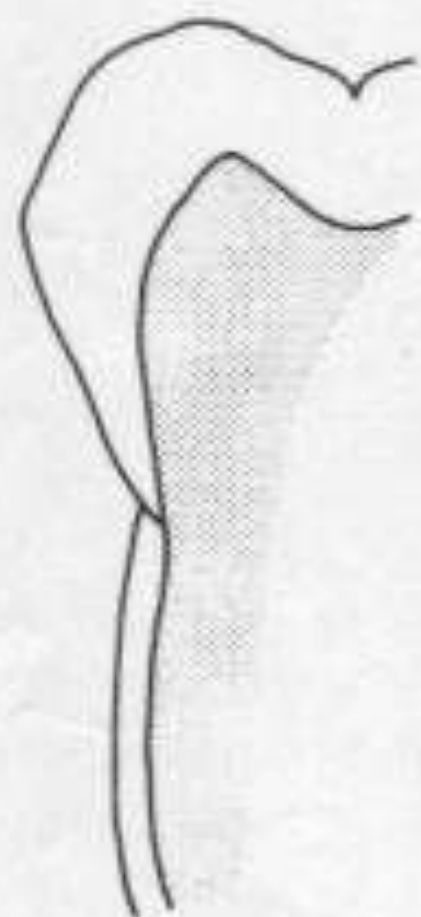
Acellular cementum (AC) showing incremental lines running parallel to the long axis of the tooth. These lines represent the appositional growth of cementum. Note the thin, light lines running into the cementum perpendicular to the surface; these represent Sharpey's fibers of the periodontal ligament (PL). D, Dentin.

Cemento-enamel junction(C.E.J):

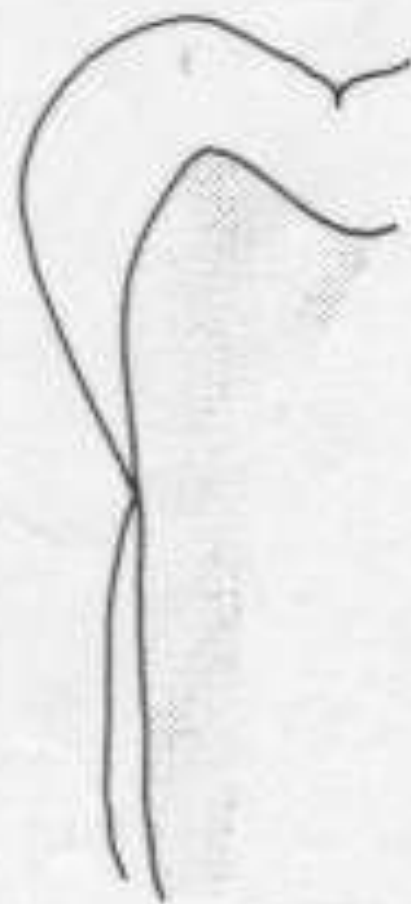
There are three types of relationships involving the cementum may exist at the C.E.J:-

- Cementum overlaps the enamel(60-65%).**
- Edge-to edge(30%)**
- cementum and enamel fail to meet (5-10%).**

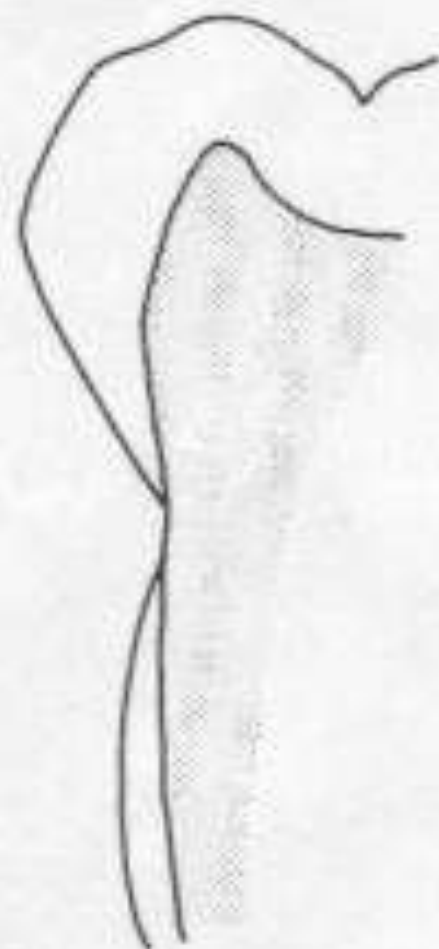
+ in the last condition, there is a possibility of gingival recession which may result in sensitivity because the dentin is exposed



Overlap
(60%-65%)



Butt
(30%)



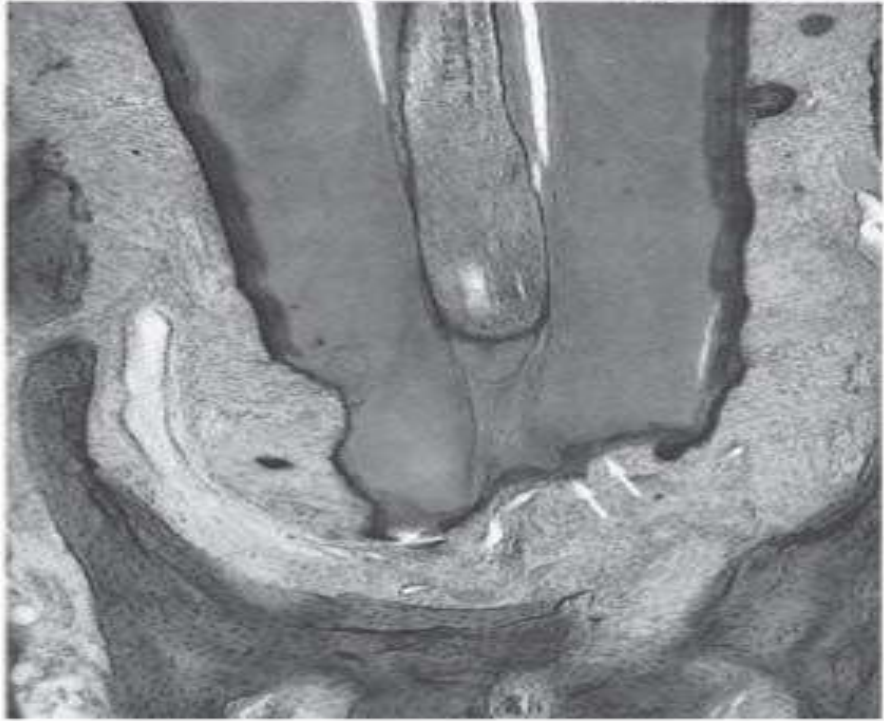
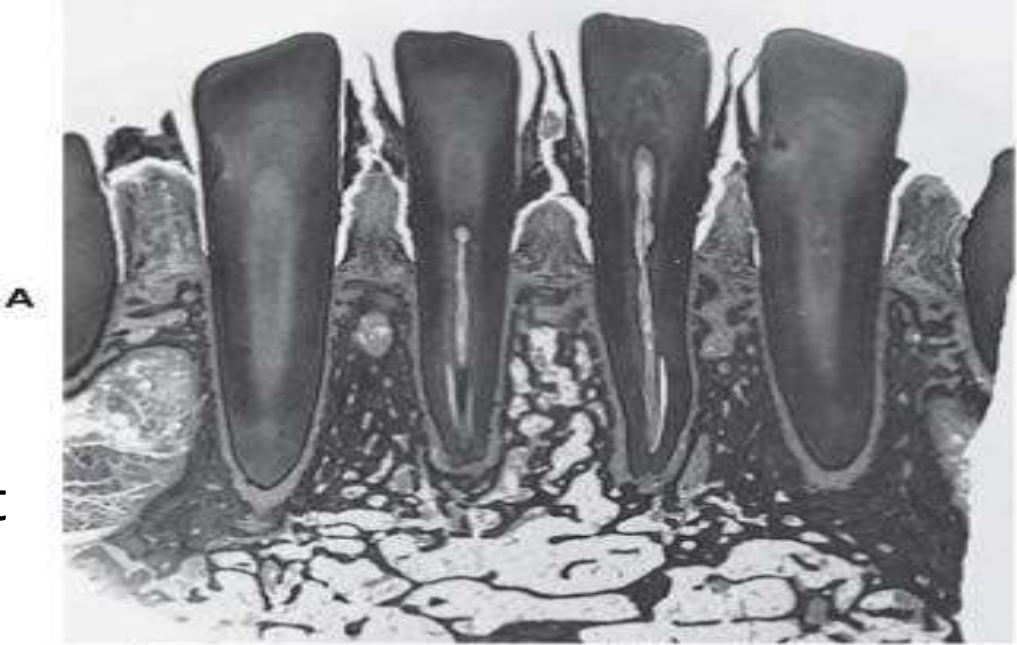
Exposed Dentin
(5%-10%)

Fig. 1-10

Cementum Resorption and Repair

- ?** Cementum resorption may be due to local or systemic causes (trauma from occlusion; orthodontic movement; cysts, and tumors; replanted and transplanted teeth calcium deficiency, hypothyroidism, Paget's disease).
- ?** Cementum resorption is not continuous, may alternate with periods of repair

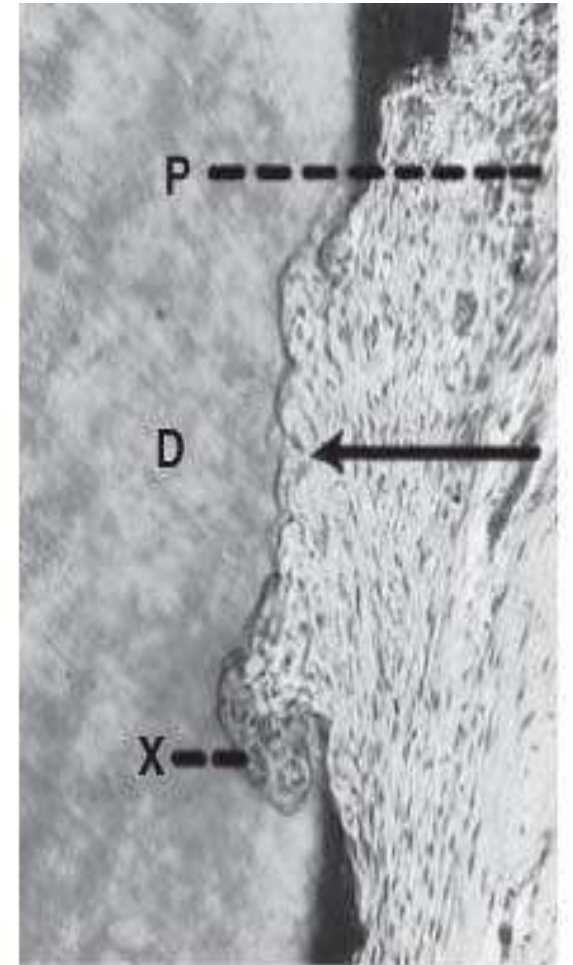
Cemental resorption associated with excessive occlusal forces. A, Low-power histologic section of mandibular anterior teeth. B, High-power micrograph of apex of left central incisor shortened by resorption of cementum and dentin. Note partial repair of the eroded areas (arrows) and cementicle at upper right.



Reversal line:

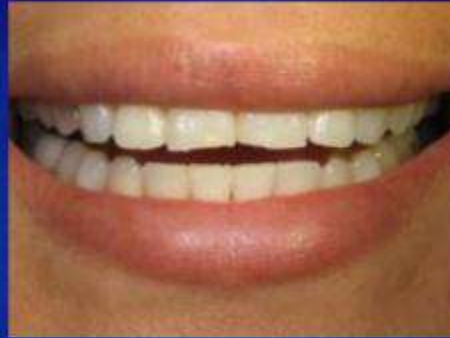
The newly formed cementum is demarcated from the root by deeply staining irregular line which delineated the border of the previous resorption.

Resorption of cementum and dentin. A multinuclear osteoclast is seen at X. The direction of resorption is indicated by the arrow. Note the scalloped resorption front in the dentin (D). The cementum is the darkly stained band at the upper and lower right. P, Periodontal ligament.



Trauma from occlusion:

Forces that exceed the adaptive capacity of the •
periodontium and produce injury.



Chronic trauma from occlusion

- is more common than the acute form and is of greater clinical significance.
- It most often develops from gradual changes in occlusion produced by **tooth wear, drifting movement, and extrusion of teeth, combined with parafunctional habits** such as bruxism and clenching,



(CARRANZA'S clinical periodontology 9th edition - 396)

Hypercementosis:

Refer to a prominent thickening of the cementum, it may be localized to one tooth e.g. tooth without antagonists or with periapical lesion, and sometimes affect the entire dentition that may occur in patient with pagets diseases.

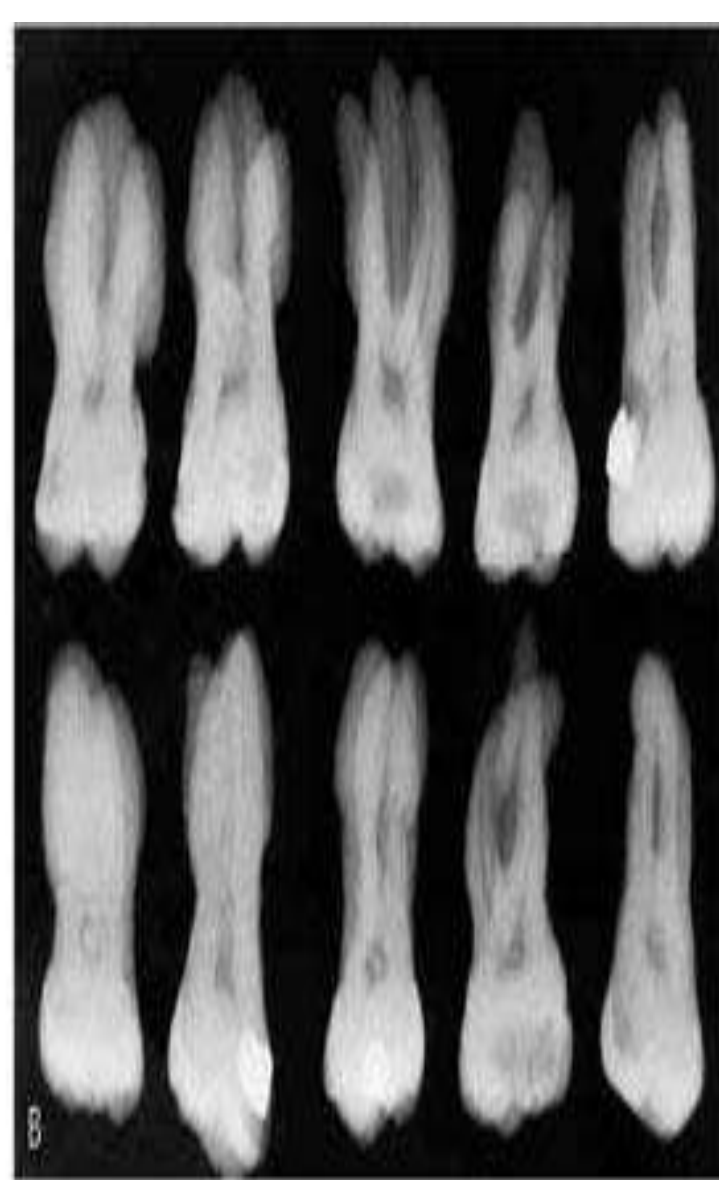


FIGURE 2 - Upper molars are the most affected teeth with hypercementosis, in various intensity levels, as these specimen, obtained from a sample of 21,573 teeth.⁶

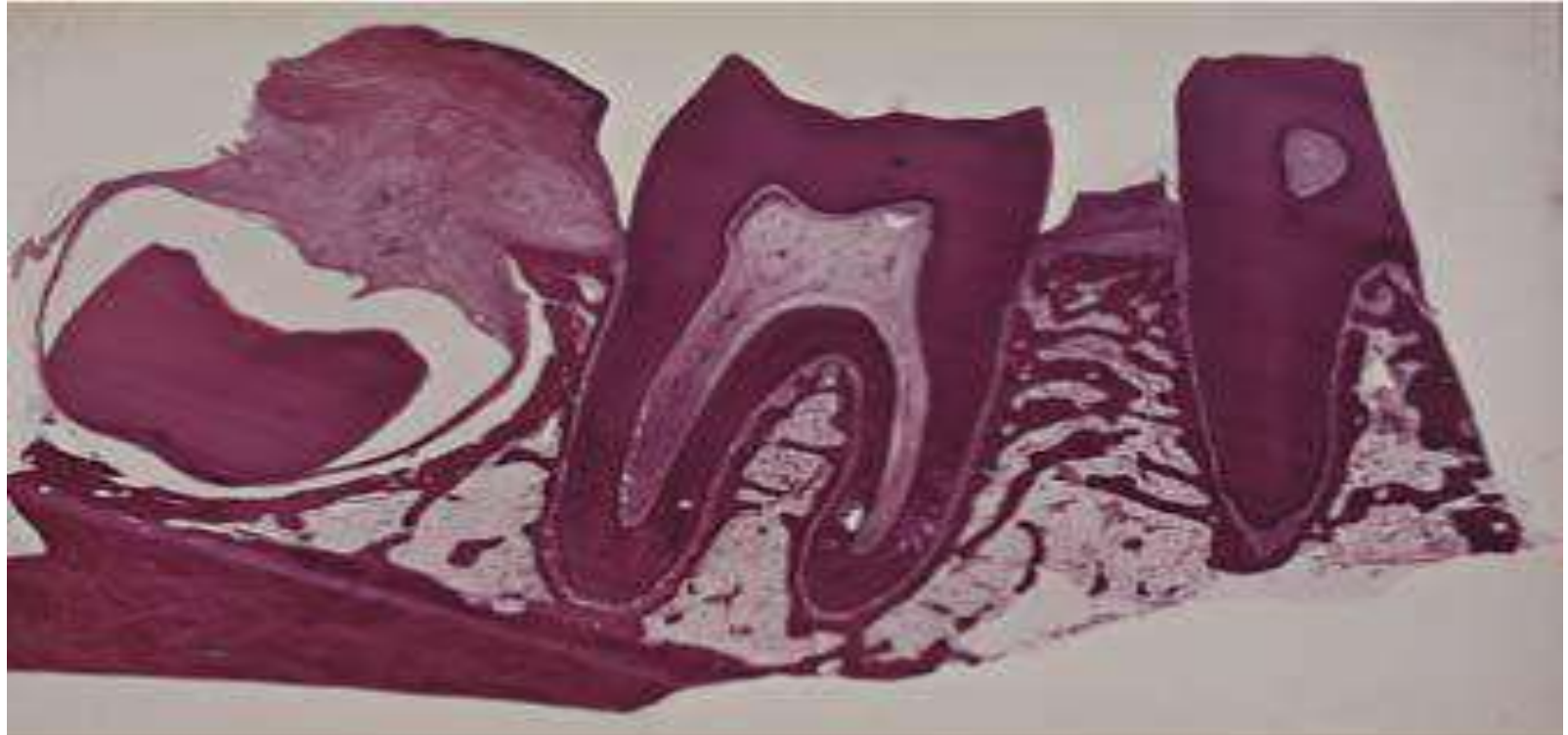
Ankylosis:

Fusion of the cementum and alveolar bone with obliteration of the PDL . It result in resorption of the cementum and it gradual replacement by bone tissue and it may develop after chronic periapical inflammation and occlusal trauma.

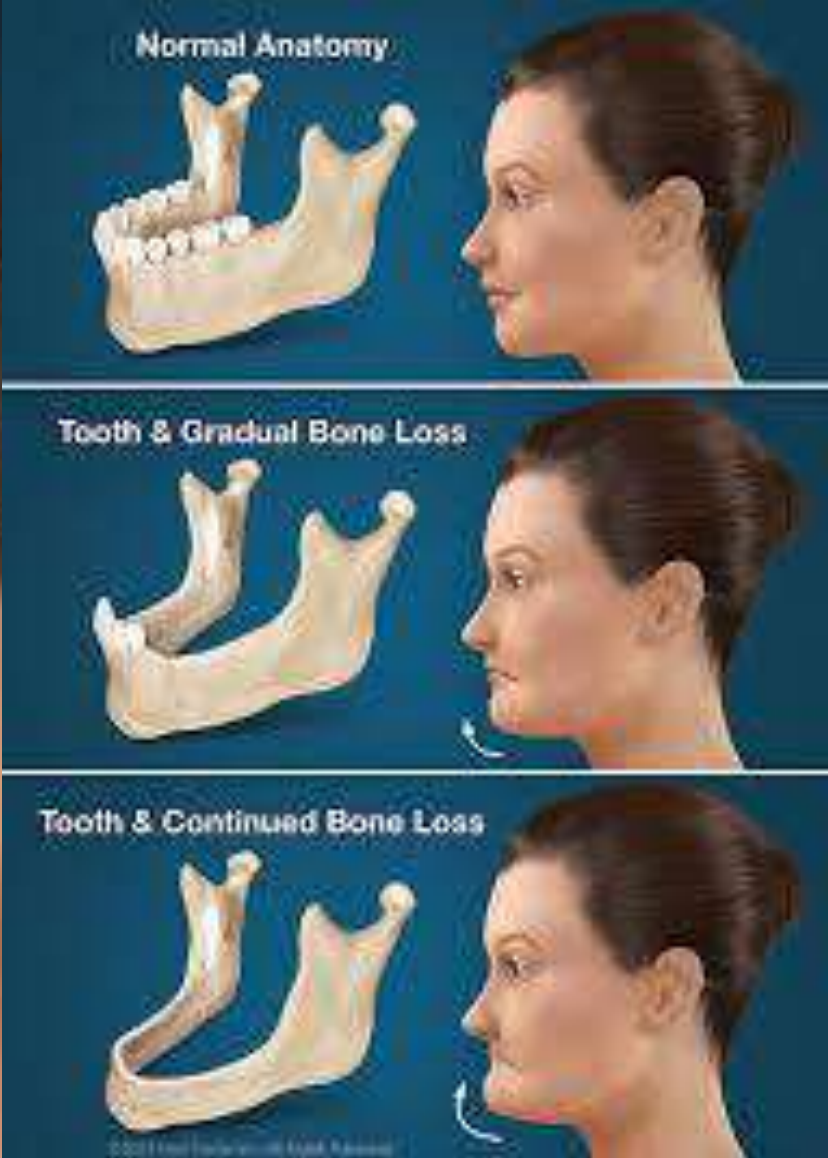
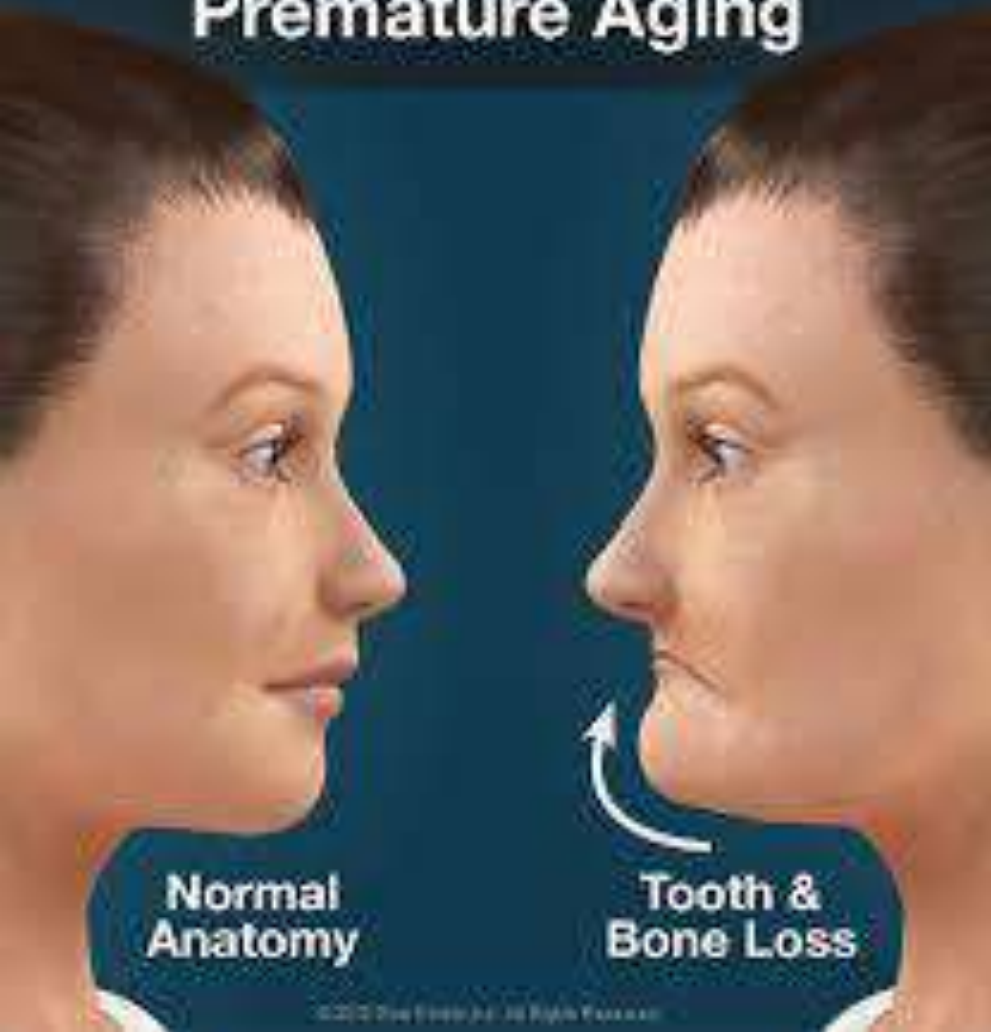


Alveolar process(AP):

- Is the portion of the maxilla and mandible that forms and support the tooth socket(alveoli).
- It develop in conjunction with the formation of and during the eruption of the teeth and it is gradually resorbed if the teeth are lost, thus it is tooth dependent structure.



Tooth Loss & Premature Aging



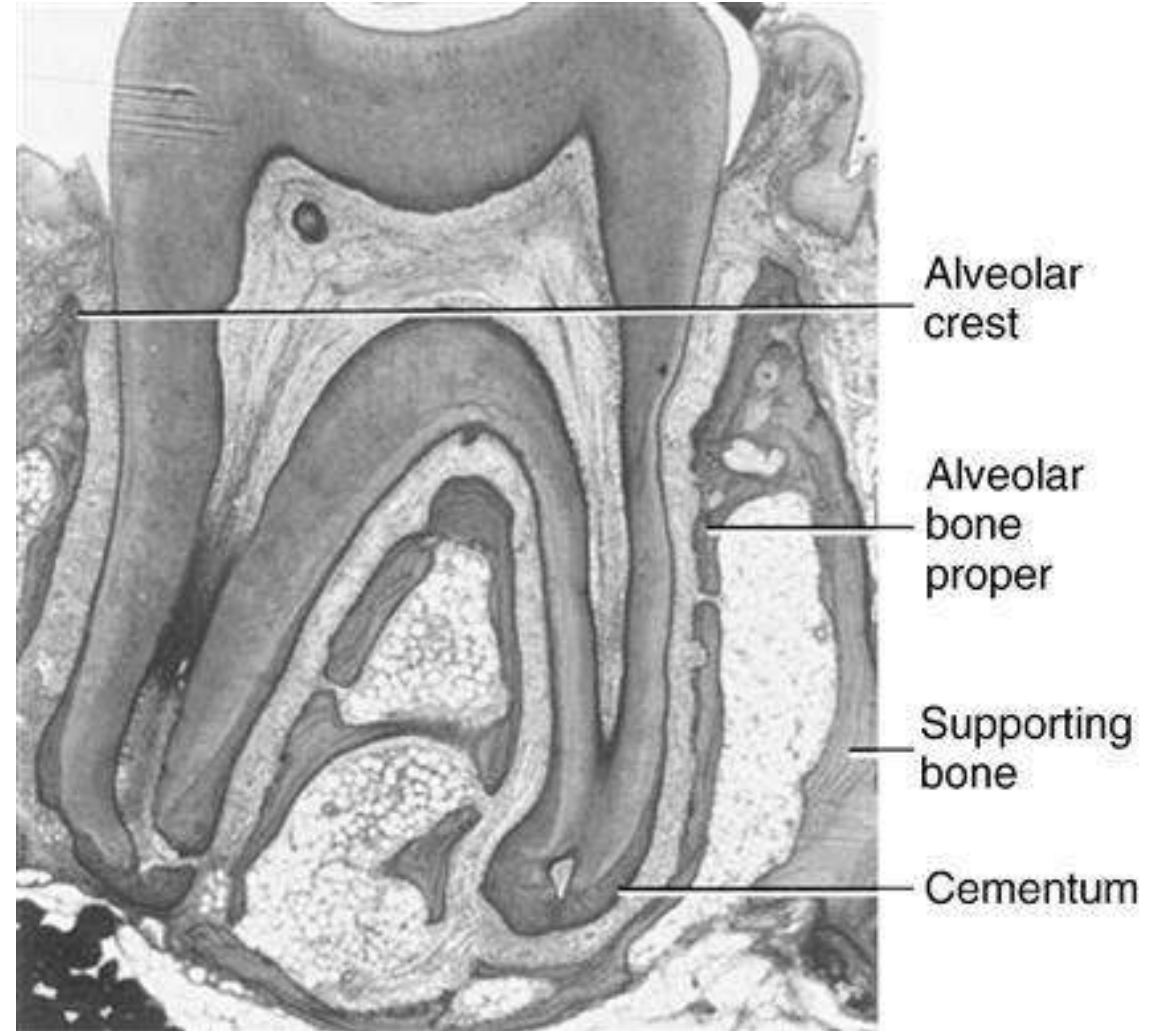
Functions of alveolar process:

- 1-comprises the attachment apparatus and the supporting tissue of the teeth together with root cementum and PDL fibers.
- 2-provide the osseous attachment to PDL fibers.
- 3- Distribution and resorb forces generated by mastication and other tooth contact.

Alveolus: is the space in the alveolar bone that accommodates the roots of the teeth.

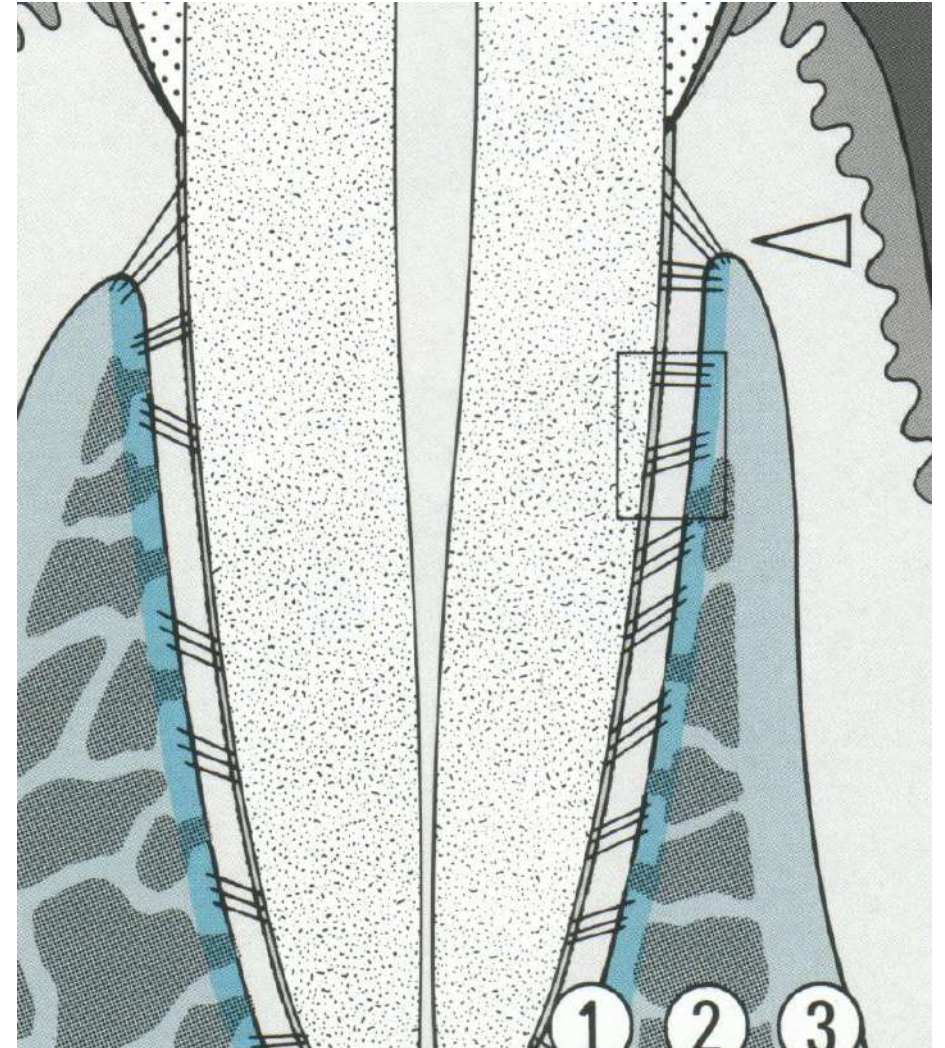
Parts of the alveolar process:

- 1- Alveolar bone proper (bundle bone) •
- 2- An external plate of cortical bone •
- 3- cancellous trabeculae or spongy bone •



Lamina Dura:

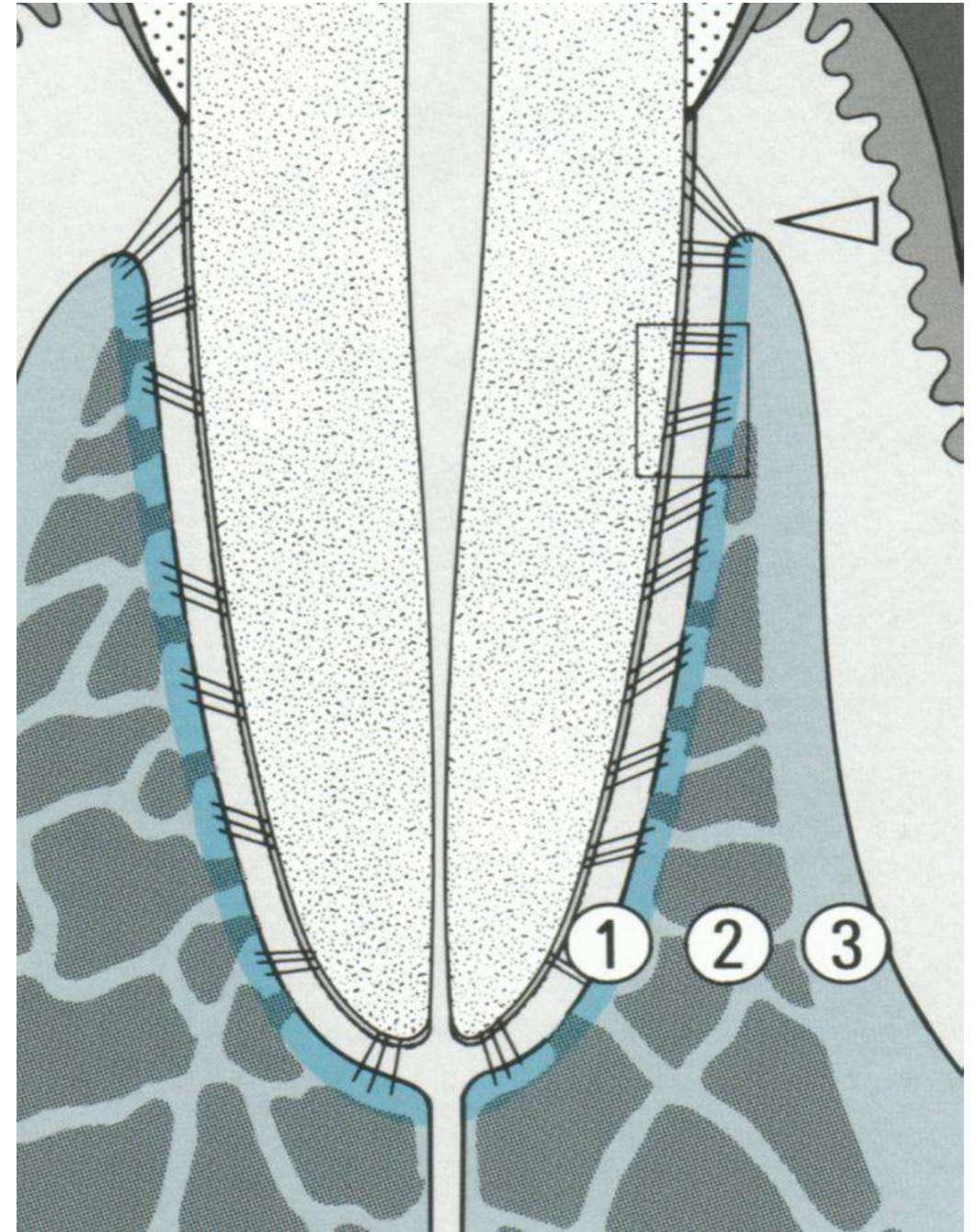
The layer of alveolar bone proper (ABP) appears a white line surrounding the root of the tooth on radiographs.

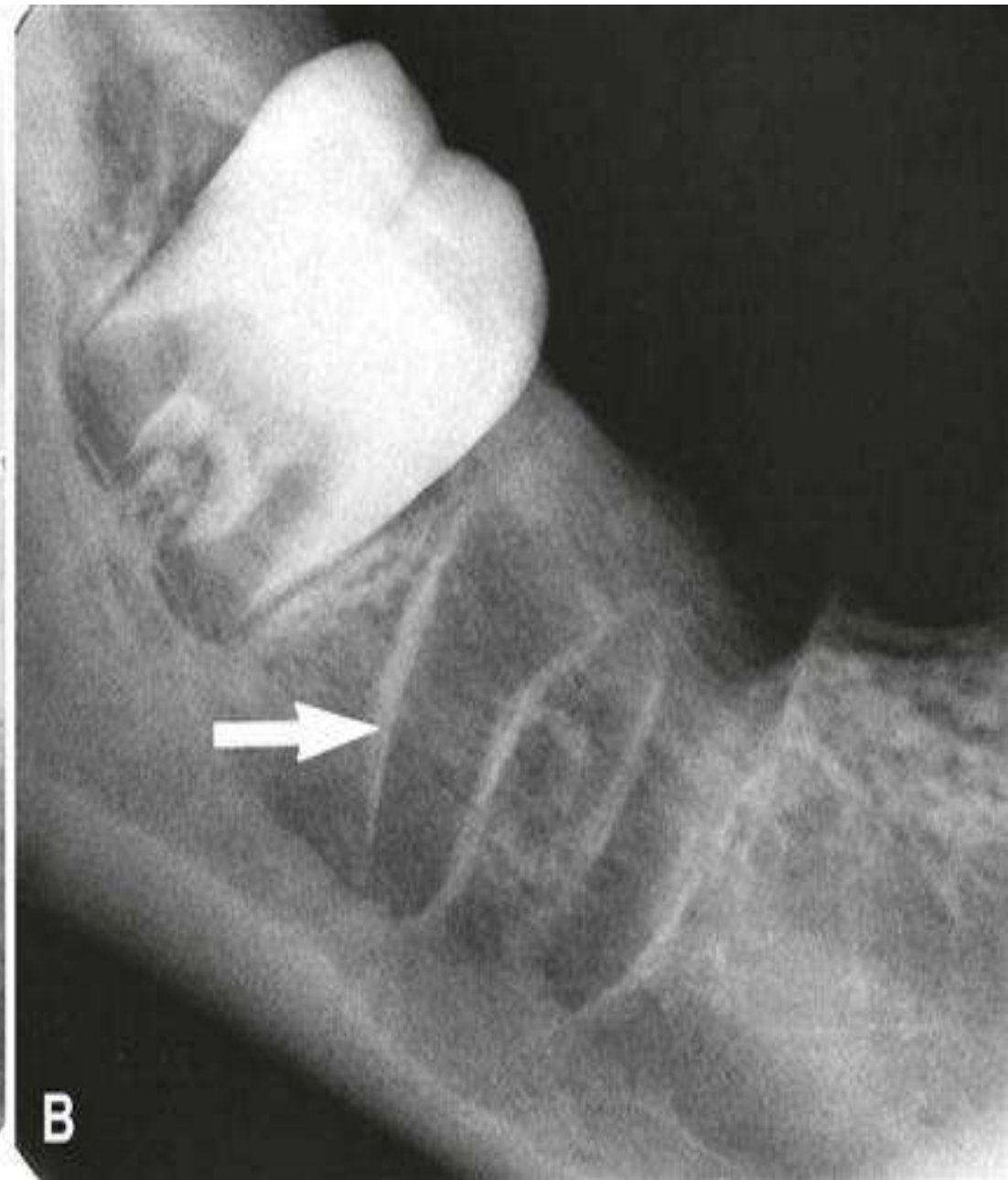


- 1 Alveolar bone, or
Cribiform plate?
Alveolar wall?
Lamina dura?**

- 2 Trabecular bone**

- 3 Compact bone**





The alveolar process are subdivided according to their anatomical relationship to the teeth :

1-Interproximal bone (interdental septum)(between two teeth)

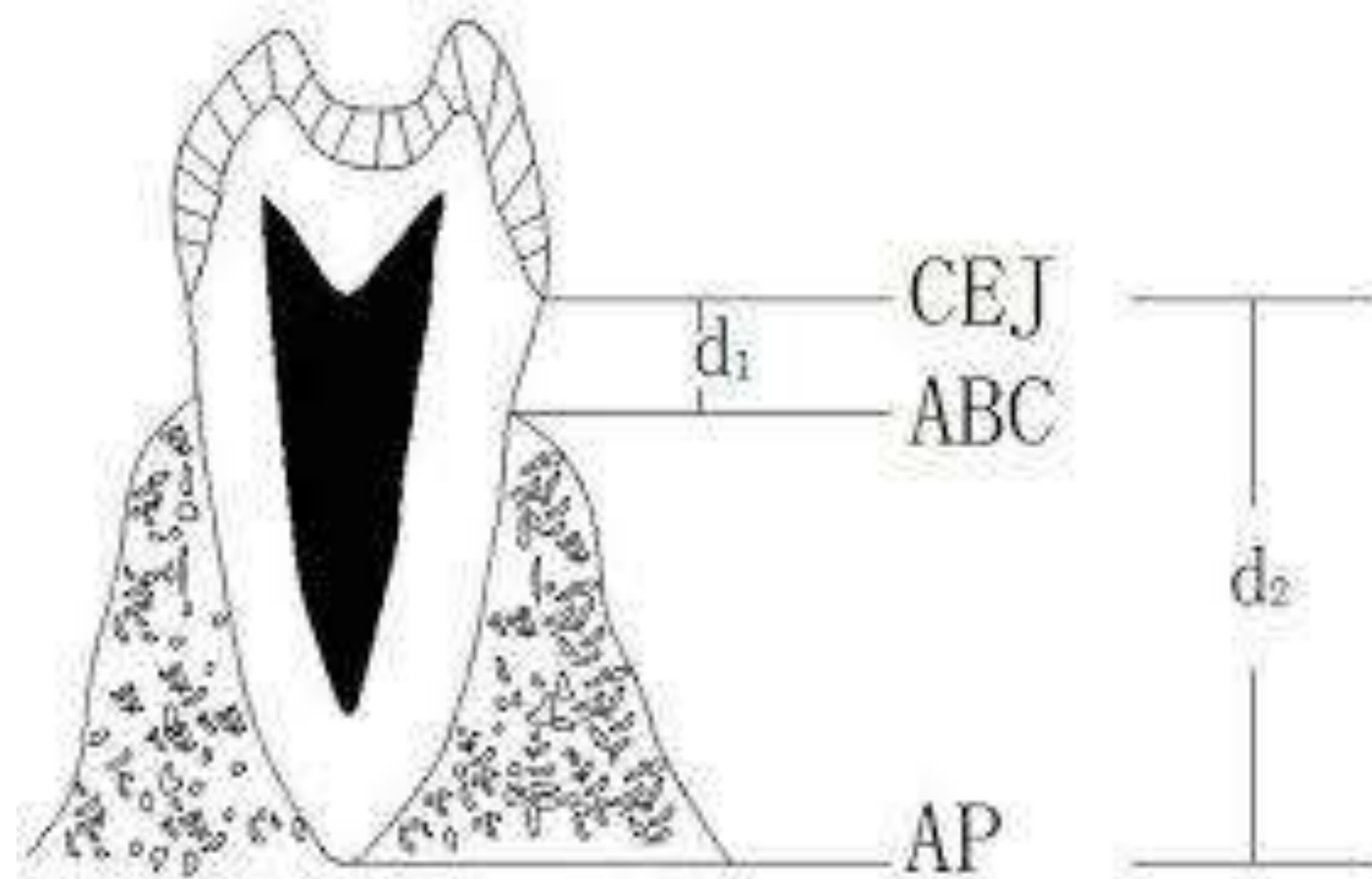
2-Inter radicular bone.(between two root in multirooted teeth_

3-Radicular bone.(located in facial, palatal, lingual surface of teeth)

 **The distance between crest of alveolar bone and C.E.J is about 2.81mm.**

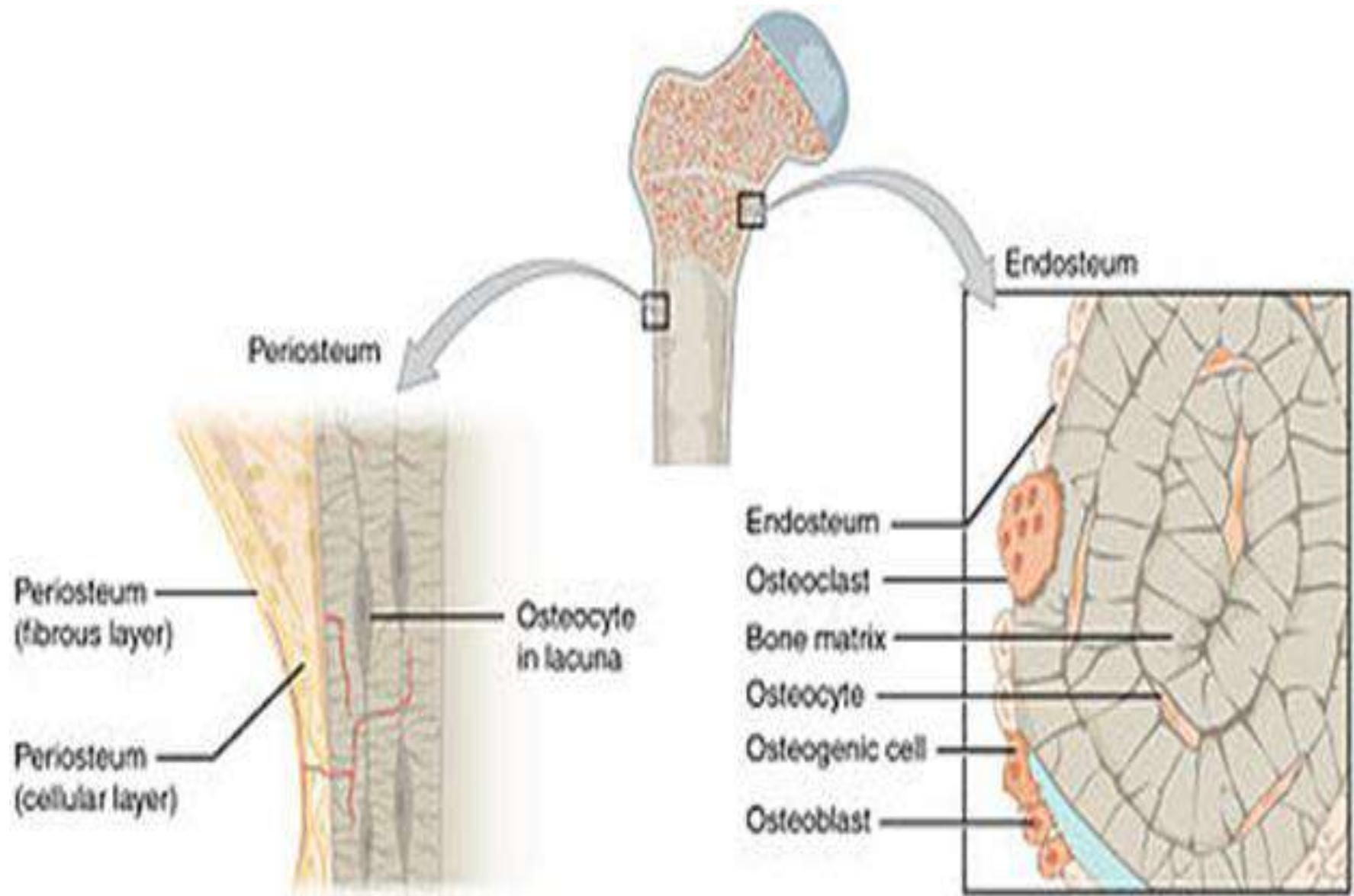


Figure 1- Focal chronic sclerosing osteitis related to pulp necrosis and chronic periapical lesion, with small and irregular thin bone areas around the root canal opening, with predominant radiopaque images in the surroundings of thin bone areas.



Periosteum: it is a layer of tissue covering the outer surface of bone, it contains collagen fibers and cells (osteoblast) with blood vessels, nerves and fibers.

Endosteum: the marrow spaces inside the bone are lined by •
endosteum, this tissue contains cells (osteoblasts).



Bone cells:

1-Osteoblast cells(bone forming cells).

2- osteoclasts cells.

3-osteocyte cells.



Composition of the bone:

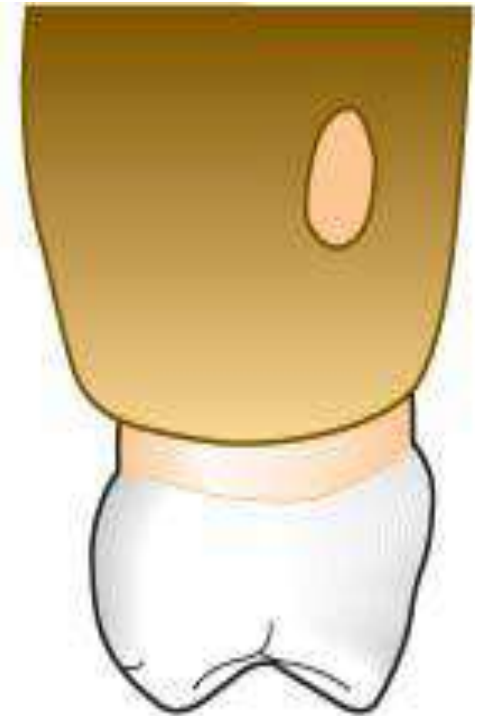
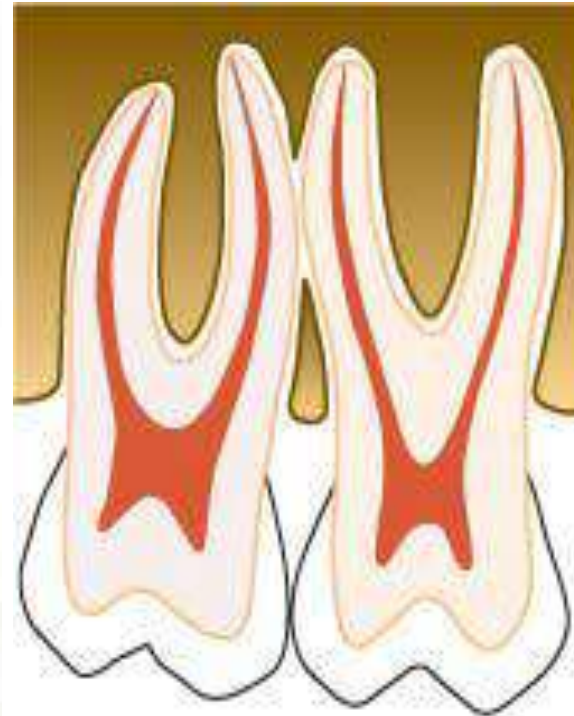
1- consist of **2/3 inorganic** matter (minerals < • calcium and phosphate >)

2- **1/3 organic** matter consist of 90% of collagen • fibers

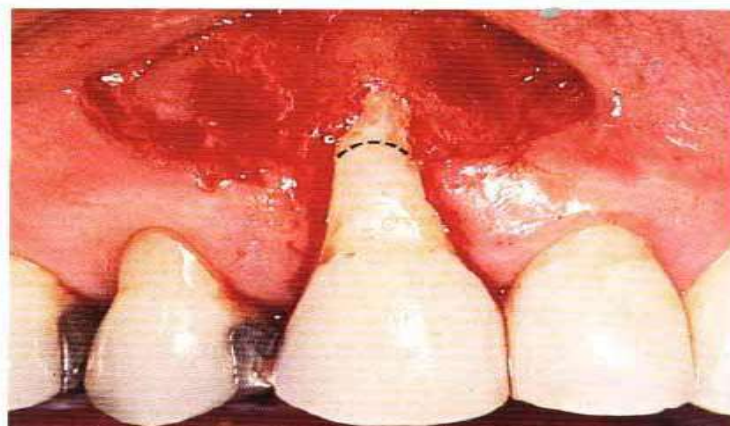
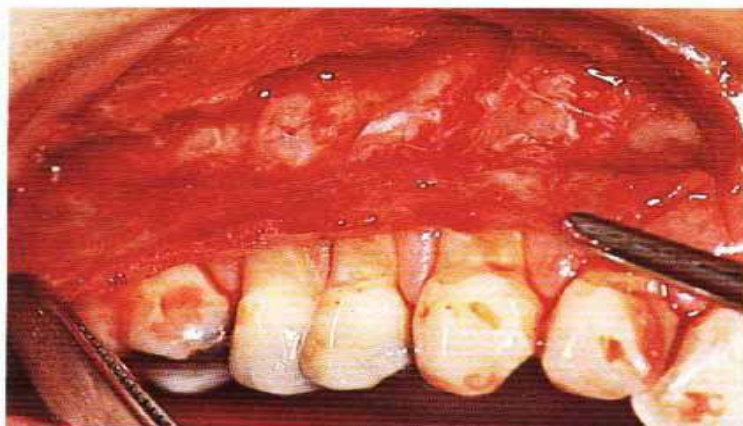
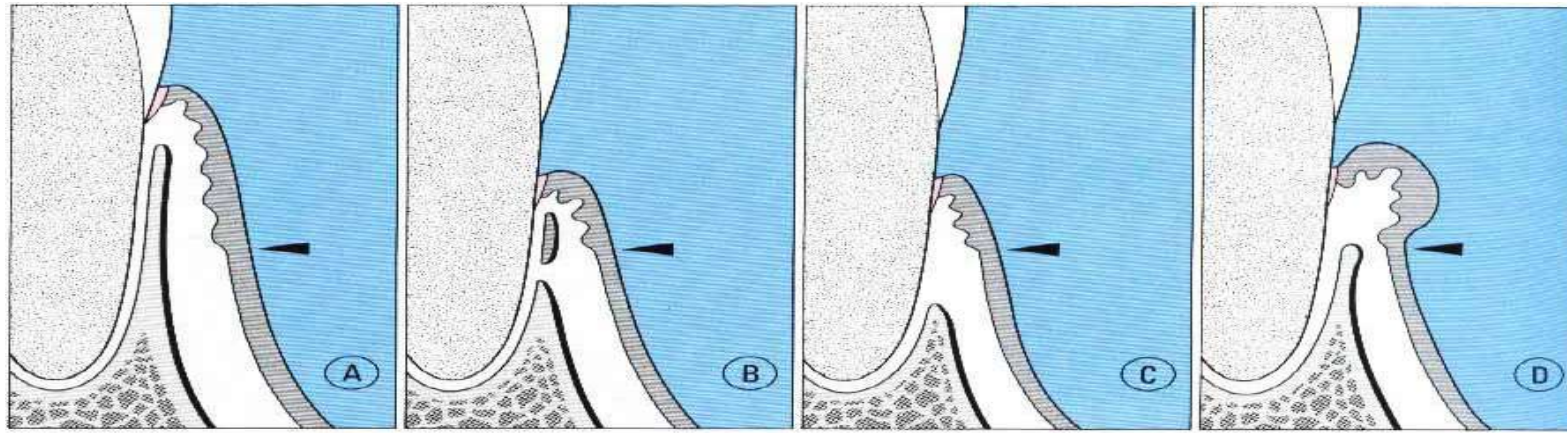
Anatomical defects of bone:

1- Fenestration (Window)

2- Dehiscence



Copyright © 2006 by Saunders, an imprint of Elsevier Inc.



Remodeling of Alveolar Bone

- Internal remodeling (resorption and formation), which are regulated by local and systemic influences.
- Local influences include functional requirements on the tooth as well as age-related changes in bone cells.
- Systemic influences are probably hormonal (parathyroid hormone, calcitonin, and others).
- Remodeling of alveolar bone affects its height, contour, and density

-

Physiologic Migration of the Teeth

- **Tooth movement does not end when active eruption is completed and the tooth is in functional occlusion.**
- **With time and wear, the proximal contact areas of the teeth are flattened and the teeth tend to move mesially. This is referred to as physiologic mesial migration. By age 40, it results in a reduction of about 0.5 cm in the length of the dental arch from the midline to the third molars. Alveolar bone is reconstructed in compliance with the physiologic mesial migration of the teeth.**
- **Bone resorption is increased in areas of pressure along the mesial surfaces of the teeth, and new layers of bundle bone are formed in areas of tension on the distal surface**

Comparison of cementum upon bone:

Cementum like bone in many aspect but different in:

1- Has no innervation.

2- Has no blood or lymph vessels .

3- does not undergo physiological remodeling (resorption and deposition), but it is characterized by continuous deposition throughout life.

4-It is a microscopic organization.


بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ



Chemical plaque control

اعداد:

د. نور صباح ارحيم



Gingivitis and periodontitis are highly prevalent diseases and prevention of occurrence or recurrence is dependent on supra-gingival plaque control. Tooth cleaning is largely influenced by the compliance and dexterity of the individuals thus the concept of using chemical plaque control is just an adjunctive mean to overcome inadequacies of mechanical cleaning.

- The action of these chemicals could fit into four categories: -

1-Anti-adhesive. -

2-Antimicrobial. -

3-plaque removal -


4-Anti-pathogenic. -

Anti-adhesive agents

They would act at the pellicle surface to prevent the initial attachment of the primary plaque forming bacteria and development of biofilms, although the amine alcohol, delmopinol, which appears to interfere with bacteria matrix formation and therefore fits between the concepts of anti-adhesion and plaque removal. Has been shown effective against plaque and gingivitis.



Antimicrobial agents

They could inhibit plaque formation through one of two mechanisms  alone or combined. The first would be the inhibition of bacterial proliferation therefore could exert their effects either at the pellicle coated tooth surface before the primary plaque formation bacteria attach or after attachment but before division of these bacteria, this effect would be bacteriostatic in type while, the second effect could be bactericidal, whereby the antimicrobial agents destroys all of the microorganisms either attaching or already attached to the tooth surface.





Plaque removal agents ➡

Such agents contained in a mouth rinse to reach all ➡ tooth surfaces and act in an identical manner to a tooth brush and remove bacteria from a tooth surface have attracted the terminology of the chemical tooth brush e.g. hypochlorites.



Anti-pathogenic agents

These agents might inhibits the expression of plaque 
microorganisms pathogenicity without necessarily
destroying them e,g. antimicrobial agents with
bacteriostatic effects.



Chlorhexidine digluconate (CHX) ➤

-CHX is frequently used as a mouth rinse(0.2% or 0.12%). The compound can ➤ also be applied as a gel, spray, varnishes, and has been incorporated into tooth paste, chewing gum, periodontal pack and sub gingival irrigation.

-at low concentration, CHX is bacteriostatic, at high ➤ concentration , it is bactericidal. The mode of action of CHX in killing bacteria is dependent upon the drug having access to cell walls.

This is facilitated by electrostatic forces, since CHX is positively charged , ➤ while the phosphate and carboxyle groups of bacterial cell walls carry negative charged . Binding causes disruption of the osmotic barrier and interfere with membrane transport.



-rinsing with CHX reduce the number of bacteria in saliva by between 50% and 90%. A maximum reduction of 95% occur around 5 days. After which the number increase gradually to maintain an overall reduction of 70%-80% at 40 days.

-An important property of CHX is its substantivity that is, the retention in the mouth and subsequent release from oral structures. After a 1 minute oral rinse of 10ml CHX 0.2% approximately 30% of the drug is retained, within 15 seconds of rinsing, half will be bonded to receptor molecules.

Clinical uses of CHX: ➤

- 1-As an adjunct to oral hygiene and professional prophylaxis. ➤
- 2-post oral surgery including periodontal surgery or rot planning. ➤
- 3-for patients with jaw fixation. ➤
- 4-medically compromised individuals predisposed to oral infections. ➤
- 5-high risk caries patients. ➤
- 6-in denture stomatitis. ➤
- 7-oral mal Oдер. ➤
- 8-recurrent oral ulceration. ➤
- 9-removable and fixed orthodontic appliance wearer. ➤
- 10-immediate preoperative CHX rinsing and irrigation. ➤
- 11-reduce salivary flow. ➤
- 12-for oral and gingival health benefits in mentally and physically handicapped. ➤

Side effects of CHX: ➡

- 1-Brown discoloration of the teeth and some restoration materials ➡ and the dorsum of the tongue.
- 2-Taste perturbation where the salt taste appears to be preferentially ➡ affected to leave food and drinks with a rather bland taste.
- 3-Enhanced supra-gingival calculus formation. ➡
- 4-Oral mucosal erosion. ➡
- 5-unilateral or bilateral parotid swelling. ➡
- 6-CHX also has a bitter taste which is difficult to mask completely. ➡



-CHX is non toxic even if digested or topically applied and has a broad antimicrobial action including wide range of gram positive and gram negative m.o., it is also effective against fungi and yeast including candida some viruses (HIV and HBV).

-It was demonstrated that rinsing for 60 seconds twice per day with 10ml of 0.2% gluconate solution in absence of tooth cleaning inhibited plaque regrowth and development of gingivitis. After that the patients should not eat or drink anything for up to 30min. With tooth brushing by using tooth paste, CHX mouth wash should be used after 30min. Of brushing ;otherwise cross reaction may occur and reduce the plaque inhibition of CHX.

Studies suggest a slow release of CHX from surface to produce a persistence bacteriostatic action lasting for about 12hr. That's why it should be used twice a day.



THANK



YOU

Classification of periodontal diseases and conditions (2017)

م. سہا اسود دہش

B.D.S, MSc. Periodontology

ATTEMPTS AT CLASSIFICATION:

- Classification of disease is necessary to try to separate conditions into distinct categories so as to aid clinical and laboratory diagnosis and specific treatment.
- The criteria for separating diseases in this way should ideally be based on etiology, histopathology and, where appropriate, genetics rather than age of onset and rates of disease progression. Over the last three decades there have been **four major attempts** to classify periodontal disease.
- Major changes were made in the **1999 classification of periodontitis**, which has been in use for the last 19 years. Periodontitis was reclassified as **chronic, aggressive (localized and generalized), necrotizing and as a manifestation of systemic disease**.

- The workshop in 2017 agreed on a classification framework for periodontitis further characterized based on a multidimensional staging and grading system that could be adapted over time as new evidence emerges

Classification of periodontal diseases and conditions (2017):

- **1- Periodontal health and gingival diseases and conditions**
 - Periodontal health and gingival health
 - Dental biofilm induced gingivitis
 - Non-dental biofilm induced gingival disease
- **2- Periodontitis**
 - Periodontitis
 - Necrotizing periodontal diseases
 - Periodontitis as a manifestation of systemic disease
- **3- Other conditions affecting the periodontium**
 - Periodontal abscess and endodontic periodontal lesions
 - Mucogingival deformity and conditions
 - Traumatic occlusal force
 - Tooth and prosthetic related factors

- **4- Peri-implant disease and conditions**

- **Peri- implant health**
- **Peri-implant mucositis**
- **Peri-implantitis**
- **Peri-implant soft and hard tissues deficiency**

1. Periodontal health and gingival diseases and conditions

- **Periodontal health and gingival health:**
- absence of clinically detectable inflammation.



Image 1

Pristine periodontal health, a very rare condition



Clinical gingival health can be restored following treatment of gingivitis and periodontitis.

bleeding on probing score (BOP%)

- assessed as the proportion of bleeding sites when stimulated by a standardized (dimensions and shape) periodontal probe with a controlled (~ 0.25 N) force to the apical end of the sulcus.



Clinical gingival health on an intact periodontium

- Absence (or minimum) bleeding on probing (less than 10%).
- Absence of attachment and bone loss.
- Probing depth $\leq 3\text{mm}$.



Image 2

Clinical gingival health, <10% localised sites of bleeding



Clinical gingival health on a reduced periodontium that include:

- **Non-periodontitis patient (e.g. recession, crown lengthening)**
 - absence (or minimum) bleeding on probing (less than 10%).
 - *presence of reduced clinical attachment and bone levels.*
 - probing pocket depth ≤ 3 .
- **Stable periodontitis patient:**
 - absence (or minimum) bleeding on probing (less than 10%).
 - *presence of reduced clinical attachment and bone levels.*
 - probing pocket depth ≤ 4 .
 - no bleeding on probing at site with 4mm pocket depth.



Image 3

Gingival health/stability
on a reduced periodontium
in a periodontitis patient



Periodontal Health on a Reduced Periodontium
in a Non-periodontitis Patient. This is a 54-year-
old male patient with a **history of excessive
frequent daily toothbrushing** (he brushed five
to six times a day with a hard-bristled
toothbrush).

Dental biofilm induced gingivitis

- An inflammatory lesion resulting from interactions between the dental plaque biofilm and the host's immune-inflammatory response, which remains contained within the gingiva and does not extend to the periodontal attachment (cementum, periodontal ligament and alveolar bone).



localized gingivitis(BOP score $\leq 10\%$ and $\leq 30\%$)

Generalized gingivitis(BOP score $> 30\%$)

Dental biofilm induced gingivitis

- **Gingivitis on an intact periodontium.**
- **Gingivitis on a reduced periodontium in a non-periodontitis patient (e.g., recession, crown lengthening).**
- **Gingival inflammation** on a reduced periodontium in a successfully treated periodontitis patient (remission periodontitis).

In the periodontitis patient, the term 'gingival inflammation' is used rather than 'gingivitis'

The classification of dental biofilm induced gingivitis:

- **A- Associated with bacterial dental biofilm only.**
- **B-Mediated by systemic or local risk factors**

Systemic conditions

Sex steroid hormones
Hyperglycemia
Leukemia
Smoking
Malnutrition

Oral factors enhancing plaque accumulation

- Prominent subgingival restoration margins
- Hyposalivation

Image 4

Predisposing factors
(local risk factors),
e.g. plaque retention
factors



The classification of dental biofilm induced gingivitis:

- **C- Drug-influenced gingival enlargements** Drugs that may cause gingival overgrowth include anticonvulsant (e.g. phenytoin), immunosuppressant (e.g. cyclosporine A), and calcium channel blockers (e.g. nifedipine, verapamil).

Image 5

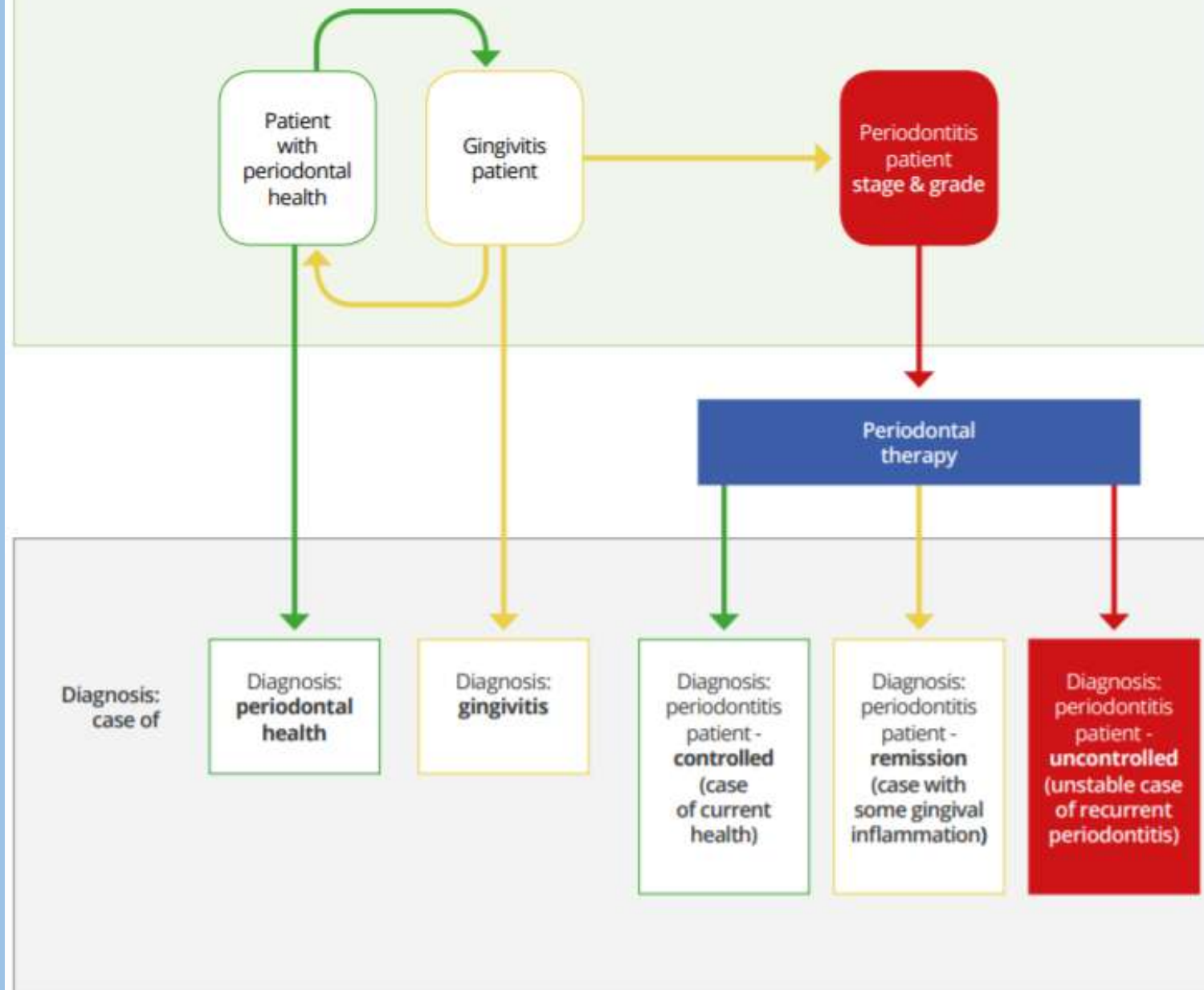
Drug influenced
gingival
enlargement



The common clinical characteristics of drug-influenced gingival enlargement include:-

- Variation in interpatient and inpatient pattern (genetic predisposition).
- Predilection for anterior gingiva.
- Higher prevalence in children and younger age group.
- Onset within 3 months of use.
- Change in the gingival contour leading to modification of gingival size.
- Enlargement first observed at the interdental papilla.
- Change in gingival color.
- Increased gingival exudate.
- Bleeding upon provocation.
- Pronounced inflammatory response of gingiva in relation to the plaque present.
- Reduction in dental plaque can limit the severity of the lesion.

Patient classification/categorisation



The classification of Non- dental biofilm induced gingival lesions:

1- Genetic/developmental disorders

Hereditary gingival fibromatosis

2- Specific infections

A- Bacterial origin

- Neisseria gonorrhoeae (gonorrhoea)
- Treponema pallidum (syphilis)
- Mycobacterium tuberculosis (tuberculosis)
- Streptococcal gingivitis (strains of streptococcus)

B- Viral origin

- Coxsackie virus (hand-foot-and-mouth disease)
- Herpes simplex 1/2 (primary or recurrent)
- Varicella-zoster virus (chicken pox or shingles affecting V nerve)

C- Fungal Candidosis

The classification of Non- dental biofilm induced gingival lesions:

3- Inflammatory and immune conditions and lesions

A- Hypersensitivity reactions

- Contact allergy
- Plasma cell gingivitis
- Erythema multiforme

B- Autoimmune diseases of skin and mucous membranes

- Pemphigus vulgaris
- Pemphigoid
- Lichen planus
- Lupus erythematosus

C. Granulomatous inflammatory conditions (orofacial granulomatosis)

- Crohn's disease
- Sarcoidosis

The classification of Non- dental biofilm induced gingival lesions:

4- Reactive processes

- Epulides

Fibrous epulis

5- Neoplasms

A- Premalignant

- Leukoplakia
- Erythroplakia

B- Malignant

- Squamous cell carcinoma
- Leukemia
- Lymphoma

6- Endocrine, nutritional, and metabolic diseases

- Vitamin deficiencies

The classification of Non- dental biofilm induced gingival lesions:

7- Traumatic lesions

A- Physical/mechanical insults

- Toothbrushing-induced gingival ulceration
- Factitious injury (self-harm)

B- Chemical (toxic) insults

- Etching
- Chlorhexidine
- Acetylsalicylic acid

C- Thermal insults

- Burns of mucosa

8- Gingival pigmentation

- Smoker's melanosis
- Amalgam tattoo

Thank you

Classification of periodontitis

م. سها أسود دهش العزاوي

B.D.S, MSc. Periodontology

In 1999 the periodontitis were classified in to:

- **Chronic periodontitis:**

- localized if $\leq 30\%$.
- generalized if $> 30\%$.
- Slight = 1-2 mm CAL , moderate = 3 - 4 mm CAL, and severe = ≤ 5 mm CAL.

- **Aggressive periodontitis: (A.P.)**

- localized if $\leq 30\%$.
- generalized if $> 30\%$.
- The term aggressive periodontitis replaced the previous name early-onset periodontitis (prepubertal, juvenile periodontitis & rapidly progressive periodontitis).

clinical features of chronic periodontitis

- Most prevalent in adults, but can occur in children and adolescents;
- Amount of destruction is consistent with the presence of local factors;
- Subgingival calculus is a frequent finding;
- Associated with variable microbial pattern;
- Slow to moderate rate of progression, but may have periods of rapid progression;
- Can be associated with local predisposing factors (e.g. tooth-related or iatrogenic factors);
- May be modified by and/or associated with systemic diseases (e.g. , diabetes mellitus)
- Can be modified by factors other than systemic diseases such as cigarette smoking and emotional stress.

clinical features of aggressive periodontitis

- Except for the presence of periodontitis, patients are otherwise clinically healthy;
- Rapid attachment loss and bone destruction;
- Familial aggregation;
- Amounts of microbial deposits are inconsistent with the severity of periodontal tissue destruction;
- Elevated proportion of *aggregatibacter actinomycetemcomitans* and, in some populations, *porphyromonas gingivalis*, may be elevated;
- Phagocyte abnormalities
- Progression of attachment loss and bone loss may be self-arresting.

- Recently, based on pathophysiology, three clearly different forms of periodontitis have been identified according to new classification system proposed by the American Academy of Periodontology (AAP) and the European Federation of Periodontology (EFP) in 2017:

1. Periodontitis.

2. Periodontitis as a direct manifestation of systemic diseases

3. Necrotizing periodontitis

Periodontal diseases and conditions

Periodontal health, gingival diseases and conditions

Periodontal and gingival health

Gingivitis: biofilm induced

Gingival diseases: non biofilm induced

periodontitis

Necrotizing periodontal diseases

Periodontitis as manifestation of systemic disease

Periodontitis

Other conditions affecting the periodontium

Systemic diseases or conditions affecting the p.l supporting tissues

P.L abscesses & endodontic p.l lesions

Mucogingival deformities & conditions

Traumatic occlusal forces

Tooth and prosthetic related factors

Peri-implant diseases and conditions

Peri-implant health

Peri-implant mucositis

Periimplantitis

Peri-implant soft and hard tissue deficiencies

Key elements in the new classification of periodontitis

- **Severity:** degree of periodontal breakdown.
- **Extent:** number and distribution of teeth with detectable breakdown.
- **Complexity of management:**
 - Probing depths.
 - Type of bone loss.
 - Furcation lesions.
 - Tooth mobility.
 - Missing teeth.
- **Rate of progression.**
- **Risk factors:** smoker, diabetes

Staging and Grading system

I. Periodontitis: was classified according to different form of staging and grading. Staging relies on the standard dimensions of severity and extent of periodontitis at presentation.

The extent and distribution for each stage described as as **molar/ incisor pattern** or **localized** if the involved sites $< 30\%$ or **generalized** if the involved site $\leq 30\%$.

Severity

Guesstimate attachment loss

- *STAGE 1*: 1-2mm *4-5mm pocket*
- *STAGE 2*: 3-4mm *6-7mm*
- *STAGE 3*: >5mm *>8mm*
- *STAGE 4*: >5mm *>8mm*



Severity

Guesstimate radiographic bone loss

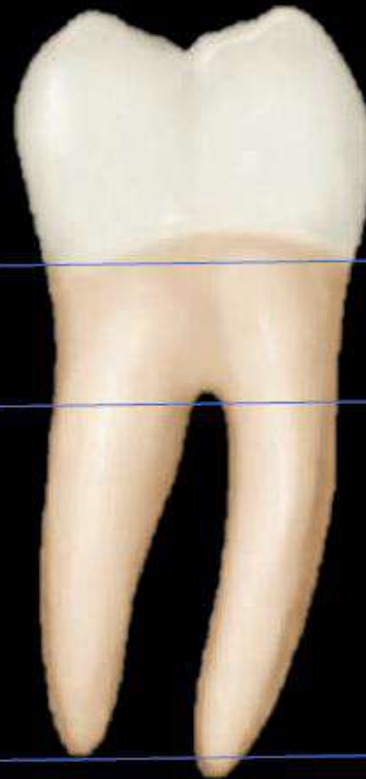


1/3

STAGE 1,2

2/3

STAGE 3,4



Staging and Grading Periodontitis



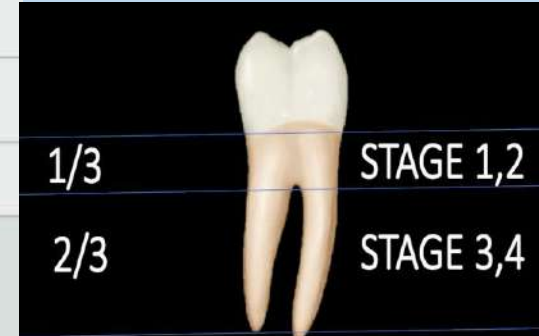
The 2017 World Workshop on the Classification of Periodontal and Peri-Implant Diseases and Conditions resulted in a new classification of periodontitis characterized by a multidimensional staging and grading system. The charts below provide an overview. Please visit perio.org/2017wwdc for the complete suite of reviews, case definition papers, and consensus reports.

PERIODONTITIS: STAGING

Staging intends to classify the severity and extent of a patient's disease based on the measurable amount of destroyed and/or damaged tissue as a result of periodontitis and to assess the specific factors that may attribute to the complexity of long-term case management.

Initial stage should be determined using clinical attachment loss (CAL). If CAL is not available, radiographic bone loss (RBL) should be used. Tooth loss due to periodontitis may modify stage definition. One or more complexity factors may shift the stage to a higher level. See perio.org/2017wwdc for additional information.

	Periodontitis	Stage I	Stage II	Stage III	Stage IV
Severity	Interdental CAL (at site of greatest loss)	1 - 2 mm	3 - 4 mm	≥5 mm	≥5 mm
	RBL	Coronal third (<15%)	Coronal third (15% - 33%)	Extending to middle third of root and beyond	Extending to middle third of root and beyond
	Tooth loss (due to periodontitis)	No tooth loss		≤4 teeth	≥5 teeth
Complexity	Local	<ul style="list-style-type: none"> Max. probing depth ≤4 mm Mostly horizontal bone loss 	<ul style="list-style-type: none"> Max. probing depth ≤5 mm Mostly horizontal bone loss 	In addition to Stage II complexity: <ul style="list-style-type: none"> Probing depths ≥6 mm Vertical bone loss ≥3 mm Furcation involvement Class II or III Moderate ridge defects 	In addition to Stage III complexity: <ul style="list-style-type: none"> Need for complex rehabilitation due to: <ul style="list-style-type: none"> Masticatory dysfunction Secondary occlusal trauma (tooth mobility degree ≥2) Severe ridge defects Bite collapse, drifting, flaring <20 remaining teeth (10 opposing pairs)
		Extent and distribution	Add to stage as descriptor	For each stage, describe extent as: <ul style="list-style-type: none"> Localized (<30% of teeth involved); Generalized; or Molar/incisor pattern 	



Staging and Grading Periodontitis



The 2017 World Workshop on the Classification of Periodontal and Peri-Implant Diseases and Conditions resulted in a new classification of periodontitis characterized by a multidimensional staging and grading system. The charts below provide an overview. Please visit perio.org/2017wwdc for the complete suite of reviews, case definition papers, and consensus reports.

PERIODONTITIS: STAGING

Staging intends to classify the severity and extent of a patient's disease based on the measurable amount of destruction of periodontal tissues as a result of periodontitis and to assess the specific factors that may attribute to the complexity of long-term case management.

Initial stage should be determined using clinical attachment loss (CAL). If CAL is not available, radiographic bone loss (RBL) should be used. Tooth loss due to periodontitis may modify stage definition. One or more complexity factors may shift the stage to a higher level. See perio.org/2017wwdc for additional information.

STAGE 3,4

	Periodontitis	Stage I	Stage II	Stage III	Stage IV
Severity	Interdental CAL <i>(at site of greatest loss)</i>	1 – 2 mm	3 – 4 mm	≥5 mm	≥5 mm
	RBL	Coronal third (<15%)	Coronal third (15% - 33%)	Extending to middle third of root and beyond	Extending to middle third of root and beyond
	Tooth loss <i>(due to periodontitis)</i>	No tooth loss		≤4 teeth	TOOTH LOSS
Complexity	Local	<ul style="list-style-type: none"> Max. probing depth ≤4 mm Mostly horizontal bone loss 	<ul style="list-style-type: none"> Max. probing depth ≤5 mm Mostly horizontal bone loss 	In addition to Stage II complexity: <ul style="list-style-type: none"> Probing depths ≥6 mm Vertical bone loss ≥3 mm Furcation involvement Class II or III Moderate ridge defects 	In addition to Stage III complexity: <ul style="list-style-type: none"> Need for complex rehabilitation due to: <ul style="list-style-type: none"> Masticatory dysfunction Secondary occlusal trauma (tooth mobility degree ≥2) Severe ridge defects Alveolar bone drifting, flaring ≥20 missing teeth (10 opposing pairs)
	Minimal complexity			HIGH COMPLEXITY	
Extent and distribution	Add to stage as descriptor	For each stage, describe extent as: <ul style="list-style-type: none"> Localized (<30% of teeth involved); Generalized; or Molar/incisor pattern 			



What is grading?

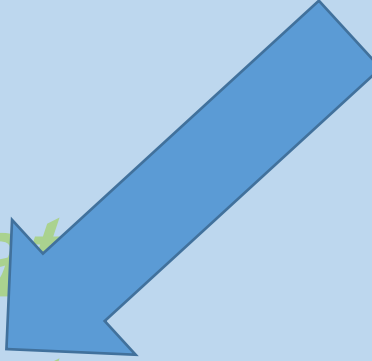
RATE
OF PROGRESSION

Grades of periodontitis:

- Irrespective of the stage at diagnosis, periodontitis may progress with different rates in individuals, may respond less predictably to treatment in some patients, and may or may not influence general health or systemic disease.
- Grading or rate of progression can be estimated by measurement of percentage of radiographical bone loss divided by the age of patient.
- **Grade A periodontitis:** is assigned if the maximum amount of radiographic bone loss in percentage terms is **less than half** the patient's age in years (for example, less than 30% in a 60-year-old or less than 40% in an 80-year-old)
- **Grade C periodontitis:** is assigned if the maximum amount of bone loss in percentage terms **exceeds the patient's age** in years (for example, more than 30% in a 28-year-old or more than 50% in a 49-year-old)
- **Grade B periodontitis:** all other situations

GRADING

START

- *GRADE A: Slow rate*
 - *GRADE B: Moderate rate*
 - *GRADE C: Rapid rate*
- 

Steps to reach diagnosis

- **Step1**: Evaluate new patient and assess whether that patient is suspect of be a periodontitis case.
- **Step2**: Determine the extent&severity.
- **Step3**: Staging.
- **Step4**:Grading.

Periodontitis as a manifestation of systemic diseases :

- 1. Systemic disorders that have a major impact on the loss of periodontal tissues by influencing periodontal inflammation:**
 - **A-** Genetic disorders.
 - **B-** Acquired immunodeficiency diseases .
 - **C-** Inflammatory diseases :
- 2. Other systemic disorders that influence the pathogenesis of periodontal diseases**
- 3. Systemic disorders that can result in loss of periodontal tissues independent of periodontitis**

Necrotizing periodontal diseases:

- Necrotizing ulcerative gingivitis
- Necrotizing ulcerative periodontitis
- Necrotizing stomatitis

Other condition affecting the periodontium:

- **A. Periodontal abscesses and endodontic periodontal lesion**

- **Periodontal abscesses (PA) :**

- **1-Periodontal abscess in periodontitis patients.**

- **A- Acute exacerbation:**

- In untreated periodontitis.
- In “refractory” periodontitis.

- **B- After different treatments:**

- Scaling and root planing or professional prophylaxis
- Surgical periodontal therapy
- Systemic antimicrobial intake, without subgingival debridement
- Use of other drugs: e.g., nifedipine.

- **2- Periodontal abscess in non- periodontitis patients.**

previously called gingival abscess



PA can also occur in previously healthy sites because of:

- Impaction of foreign bodies
- Harmful habits
- Orthodontic factors
- Gingival enlargement
- Alterations of the root surface:
 - Severe anatomic alterations, such as invaginated tooth, dens evaginatus (grooves) or odontodysplasia.
 - Minor anatomic alterations, such as cemental tears, enamel pearls or developmental grooves.
- Iatrogenic conditions, such as perforations.
- vertical root fracture or cracked tooth syndrome extending through the root.
- External root resorption.

dens evaginatus



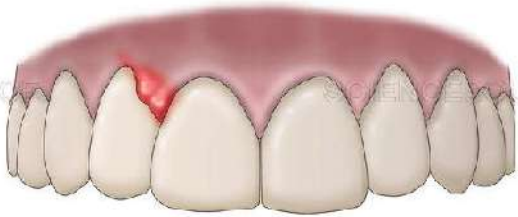
- PA may be associated with various combinations of the following clinical features:
 - Pain,
 - swelling,
 - color change,
 - tooth mobility,
 - extrusion of teeth,
 - purulence,
 - sinus tract formation,
 - fever, lymphadenopathy, and there may be a radiolucency of the affected alveolar bone.
- The acute periodontal abscess characterized by slight discomfort to severe pain and swelling. Chronic periodontal abscess is usually a symptomatic or with dull pain with a history of intermittent exudate.

The periodontal abscess need to be differentiated from the periapical abscess in the followings:

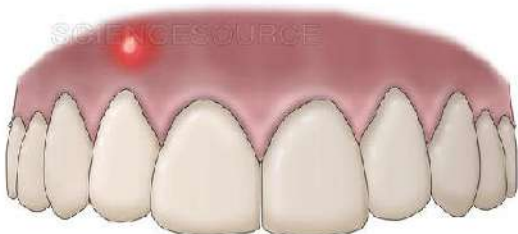
Gingival abscess



Periodontal abscess



Periapical abscess



	Periodontal abscess	Periapical abscess
1.	The tooth is vital.	Tooth is not vital.
2.	The lesion lateral to the root surface.	The lesion is most likely periapical.
3.	X-ray finding shows area of radiolucency along the lateral surface of the root.	X-ray finding shows apical radiolucency.
4.	The tooth is tender to lateral percussion.	Tooth tender to vertical percussion.

Diagnosis of Periodontal Abscess

Palpation



To check tenderness, swelling, fluctuation, and hardness in underlying tissues.

Percussion



A tooth with a periodontal abscess will be tender to percussion.

Sensitivity test



To determine if the involved tooth is vital or not.

X-ray



The periodontal abscess will appear as a dark area alongside the tooth.

Sharedentalcare
.com

Endodontic periodontal lesions:

- Clinical conditions involving both the pulp and periodontal tissues and may occur in acute or chronic forms.
- When they are associated with a recent **traumatic or iatrogenic event** (e.g. root fracture or perforation), the most common manifestation is an **abscess** accompanied by **pain**.
- endo- periodontal lesions, in subjects with **periodontitis**, normally present slow and chronic progression without evident symptoms.
- **The most common signs and symptoms :**
 - **1-** Deep periodontal pockets reaching or close to the apex.
 - **2-** Negative or altered response to pulp vitality tests.

- **The other signs and symptoms reported, in order of prevalence, are:**
 1. bone resorption in the apical or furcation region,
 2. spontaneous pain or pain on palpation and percussion,
 3. purulent exudate,
 4. tooth mobility,
 5. sinus tract,
 6. crown, and gingival color alterations

Endo-periodontal lesions associated with endodontic and periodontal infections might be triggered:

1. by a carious lesion that affects the pulp and, secondarily, affects the periodontium.
2. by periodontal destruction that secondarily affects the root canal.
3. or by both events concomitantly.

Endo-periodontal lesions associated with trauma and iatrogenic factors

- These conditions usually have a **poor prognosis** as they affect the tooth structure. The most common lesions in this category were:
 1. root/pulp chamber/furcation perforation (e.g. because of root canal instrumentation or to tooth preparation for post retained restorations)
 2. root fracture or cracking (e.g., because of trauma or tooth preparation for post-retained restorations)
 3. external root resorption (e.g., because of trauma)

B- Mucogingival deformities or conditions around teeth

- gingival biotype
 - Thin scalloped
 - Thick scalloped
 - Thick flat
- Gingival/soft tissue recession
 - Facial or lingual surfaces
 - Interproximal (papillary)
 - Severity of recession
 - Gingival thickness
 - Gingival width
- Lack of keratinized gingiva
- Decreased vestibular depth
- Aberrant frenum/muscle position
- Gingival excess
 - Pseudopocket
 - Inconsistent gingival margin
 - Excessive gingival display
 - Gingival enlargement
- Abnormal color

gingival biotype



Fig 1 *Clinical appearance of thick gingival type.*



Fig 2 *Clinical appearance of thin gingival type.*

Gingival recession

- Is location of the gingival margin apical to the cemento- enamel junction.
- The causes of gingival recession:
 - Plaque accumulation will cause destruction of the junctional epithelia as a result of the inflammatory process.
 - Traumatic gingival recession:
 - Fault tooth brushing
 - Tooth malposition
 - High frenal attachment
 - Overhanging fillings
 - Prosthetic appliances
 - Habits as nail biting.

Miller's classification of gingival recessions



C. Tooth and prosthetic related factors :

- a- Localized tooth-related factors that modify or predispose to plaque-induced gingival diseases/periodontitis
 - Tooth anatomic factors
 - Root fractures
 - Cervical root resorption, cemental tears
 - Root proximity
 - Altered passive eruption

b-Localized dental prosthesis-related factors

- Restoration margins placed within the supracrestal attached tissues
- Clinical procedures related to the fabrication of indirect restorations
- Hypersensitivity/toxicity reactions to dental materials

d.Traumatic occlusal force

- Primary occlusal trauma
- Secondary occlusal trauma
- Orthodontic force

- Occlusal trauma: Injury resulting in tissue changes within the attachment apparatus as a result of occlusal force(s).
- Primary occlusal trauma: Injury resulting in tissue changes from traumatic occlusal forces applied to tooth or teeth with normal support. It occurs in the presence of:
 - Normal bone levels, 2) Normal attachment levels, and 3) Excessive occlusal force(s).
- Secondary occlusal trauma: Injury resulting in tissue changes from normal or traumatic occlusal forces applied to a tooth or teeth with reduced support. It occurs in the presence of:
 - Bone loss, 2) Attachment loss, And 3) "Normal"/excessive occlusal force(s).

e. Peri-implant diseases and conditions

- **peri-implant health**
- In health, the peri-implant site is characterized by absence of erythema, bleeding on probing, swelling and suppuration.
- **peri-implant mucositis:** the diagnosis of peri-implant mucositis requires: Visual inspection demonstrating the presence of periimplant **signs of inflammation**: red as opposed to pink, swollen tissues as opposed to no swelling. Presence of profuse bleeding and/or suppuration on probing, an increase in probing depths compared to baseline; and absence of bone loss beyond crestal bone level changes resulting from the initial remodeling.
- **peri-implantitis:** the diagnosis of peri-implantitis will involve **radiographic** bone loss associated with gingival recession or increased probing depth in addition to signs associated with peri-implant mucositis

Thank you

Clinical
Periodontology
Fourth grade
Presented by:

Dr. NOOR SABAH IRHAYYIM

-The by-products of bacterial plaque contribute to the development of both dental caries and periodontal diseases. At-home mechanical removal of plaque is a necessary part of preventing these diseases. Regular, effective plaque removal is essential for preventing or controlling caries and periodontal diseases.

Patient's motivation:

1- Information: Aims at increasing the compliance of the patient, so the patient recognizes oral health as a valuable goal of therapy, this can be achieved by following the step by step motivation system which includes simple demonstration for:

A-Symptoms of disease vs. healthy sites, bleeding, pocket depth, recession, etc.





Healthy gingiva, explain the normal, healthy features to your patient



Inflamed gingiva, show the patient the Signs of inflammation

○ More pictures to be shown to your patient, explain to the patient what would happen if the periodontal disease left untreated (transition from health to disease):

- Healthy gingiva  Gingivitis 
periodontitis

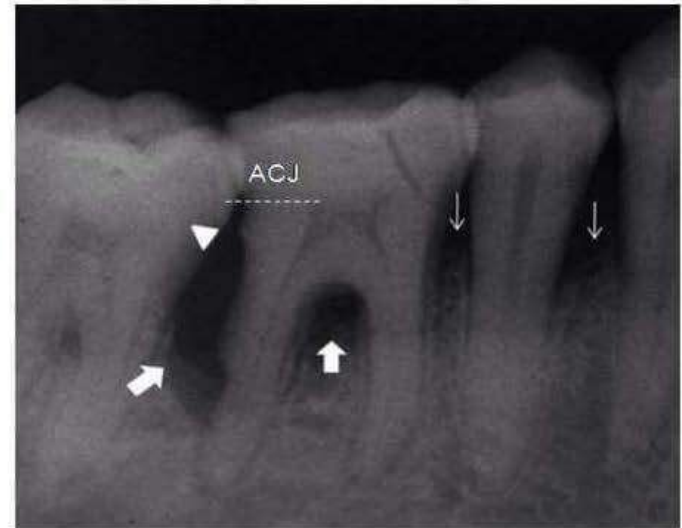
- Gingivitis (reversible), periodontitis
(irreversible) but treated to prevent further progression of the disease).

B-Radiographic bone height, normal vs. loss.



X-ray for normal periodontium

• Explain reasons of disease, bacterial plaque



Periodontitis X-ray, show the patient the bone loss (arrows)

c- Explain reasons of disease; bacterial plaque.



d- Demonstrate plaque accumulation by using **disclosing agent**, ex: erythrocin tablet, which converts the color of dental plaque into red.



Before disclosing



After disclosing
(showing plaque)

-Example of disclosing agents: disclosing solution & erythrocin tablets.



2- Testing the acquirement of knowledge: by simply asking the patient to repeat what you already demonstrate in information step, if the patient seems not understand what you have explained then repeat the information step in simpler language.

3- Acquiring the knowledge will lead to change in patient's attitude.

4- Change in patient behavior is the logical result which is expected from motivation.

Oral Hygiene Instructions

Plaque control is the regular removal of microbial plaque and the prevention of its accumulation on the teeth and adjacent gingival surfaces. Microbial plaque is the major etiology of periodontal diseases and is related to dental caries; therefore gaining patient cooperation in daily plaque removal is critical to long-term success of all periodontal and dental treatment.

Mechanical plaque control

The most common method for plaque control since it is cheap, simple and doesn't required long time to perform:

- Brushing
- Interdental tools

Chemical plaque control

It is an adjunctive to mechanical means & not consider as alternative to mechanical cleanings.

This category includes: mouth wash, dentifrices & gel.

Professional plaque control

This method is expensive, time consuming & required man power & it is indicated only in extreme cases like: severe disability, coma, etc.

Brushing method

The objectives of tooth brushing are to remove plaque, debris from the teeth and to stimulate the gingiva. There are many tooth brushing techniques the patient can use to accomplish these goals.

- Several methods suggested: Roll, Vibrating, Circular, Vertical, and Horizontal.

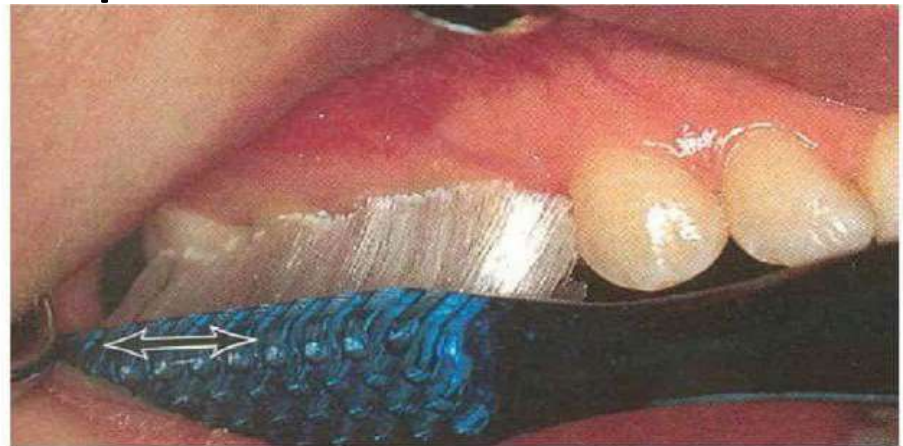
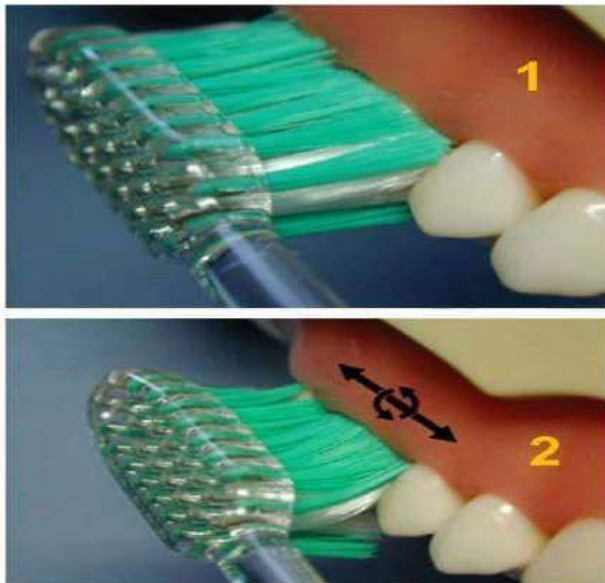
- No method is clearly superior.

-illingness and ability are more important than the technique.



1- Sulcular (Bass) method:

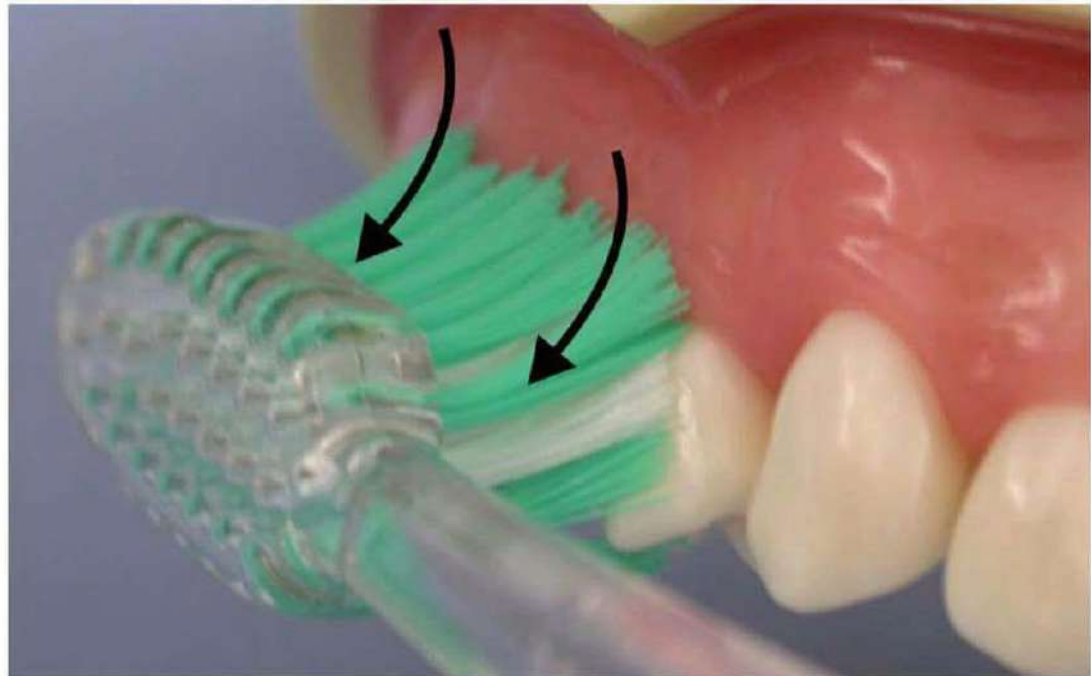
The bristles pointed at a 45-degree angle into the gingival sulcus, vibrate the brush gently forth & back about 20 times. This method is useful for patients with periodontal problems.



1. 45° angle to the gingival margin
2. Vibrate

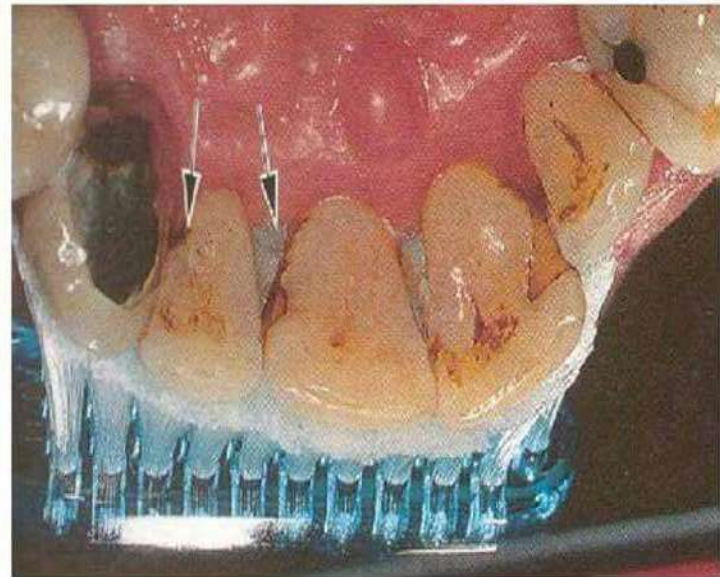
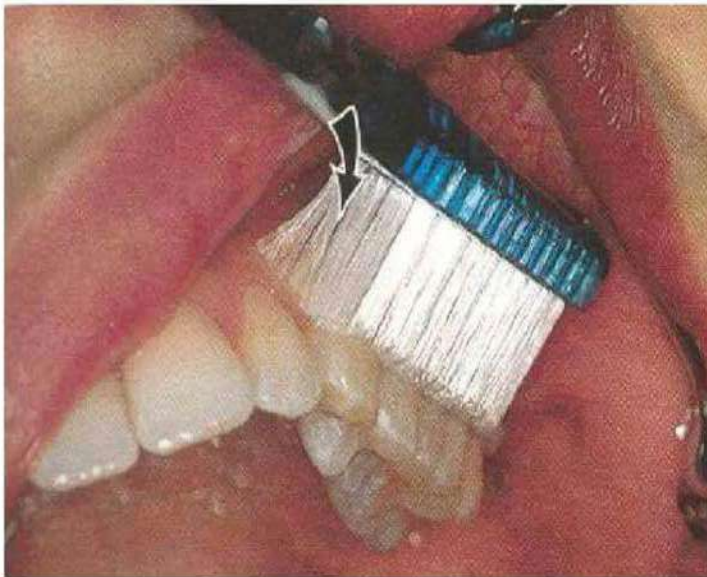
2-Rolling method:

This method is useful for stimulation of the gingiva. Place the brush above the free gingiva, exerting slight pressure; draw the brush toward the occlusal surface.



3- Charter's method:

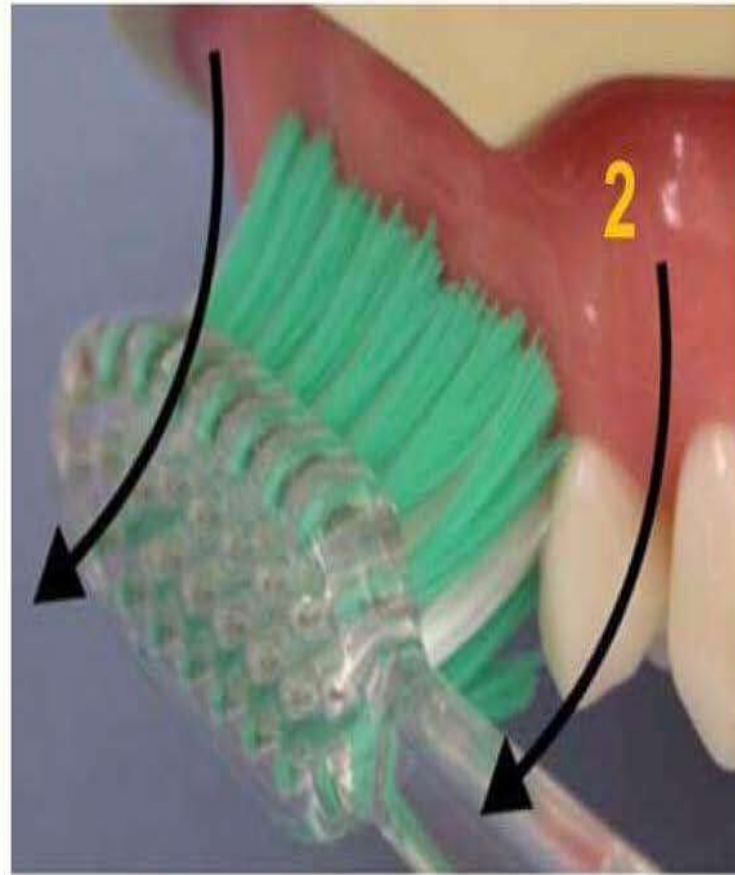
This method is useful for patients with severe loss of interdental papilla height, fixed prosthesis, previous gingival surgery, or subsided ulcerative gingivitis.



4- Modified stillman's method:

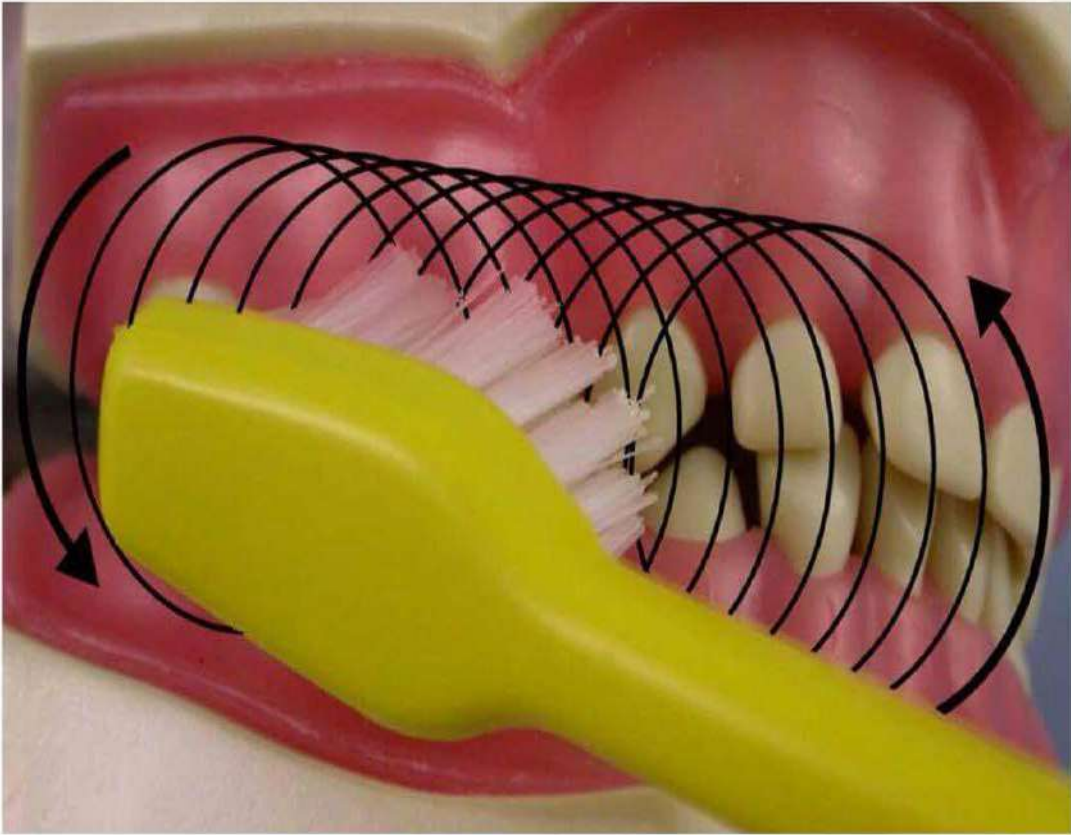


1. Press and Vibrate



2. Roll

5- fone's method:



Toothbrush

Toothbrushes vary in size and design as well as in length, hardness, and arrangement of the bristles.

However, all agree that use of hard toothbrushes, vigorous horizontal brushing, and use of extremely abrasive dentifrices may lead to cervical abrasions of teeth and recession of gingiva.

Vigorous brushing is not necessary and can lead to gingival recession, wedge-shaped defects in the cervical area of root surfaces, and painful ulceration of the gingiva.

The American Dental Association (ADA) recommends that individuals brush twice per day and use floss or other interdental cleaners once per day to effectively remove microbial plaque and prevent gingivitis. Toothbrushes must also be replaced periodically every 3 to 4 months.

Vigorous tooth brushing with an abrasive dentifrice can result in trauma to the gingiva and wearing away of the tooth surfaces, especially root surfaces, and can contribute to gingival recession.



Powered toothbrush

Powered toothbrushes have been shown to improve oral health for (1) children and adolescents, (2) children with physical or mental disabilities, (3) hospitalized patients, including older adults who need to have their teeth cleaned by caregivers, and (4) patients with fixed orthodontic appliances.

The vibrations have also been shown to interfere with bacterial adherence to oral surfaces.

INTERDENTAL CLEANING AIDS

Periodontal lesions are predominantly found in interdental locations, so tooth brushing alone is not sufficient to control gingival and periodontal diseases. It has been demonstrated in healthy subjects that plaque formation begins on the interproximal surfaces where the toothbrush does not reach.

Patients need to understand that the purpose of interdental cleaning is to remove microbial plaque, not just dislodge food wedged between teeth.

Many tools are available for interproximal cleaning and they should be recommended based on the size of interdental spaces, presence of furcations, tooth alignment, and presence of orthodontic appliances or fixed prostheses.

Also, ease of use and patient cooperation are important considerations.

Common aids are dental floss and interdental cleaners such as wooden or plastic tips and interdental brushes.

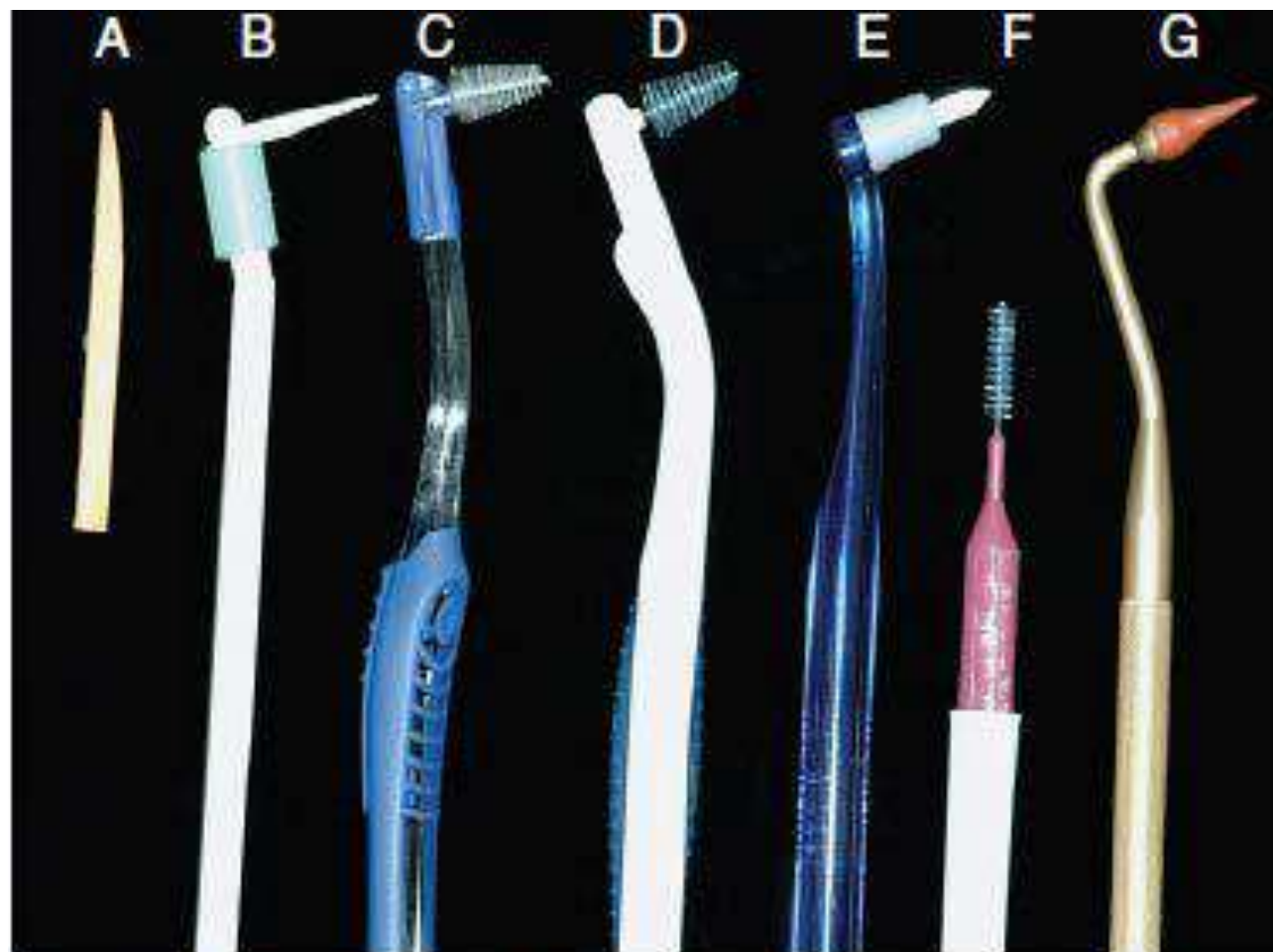


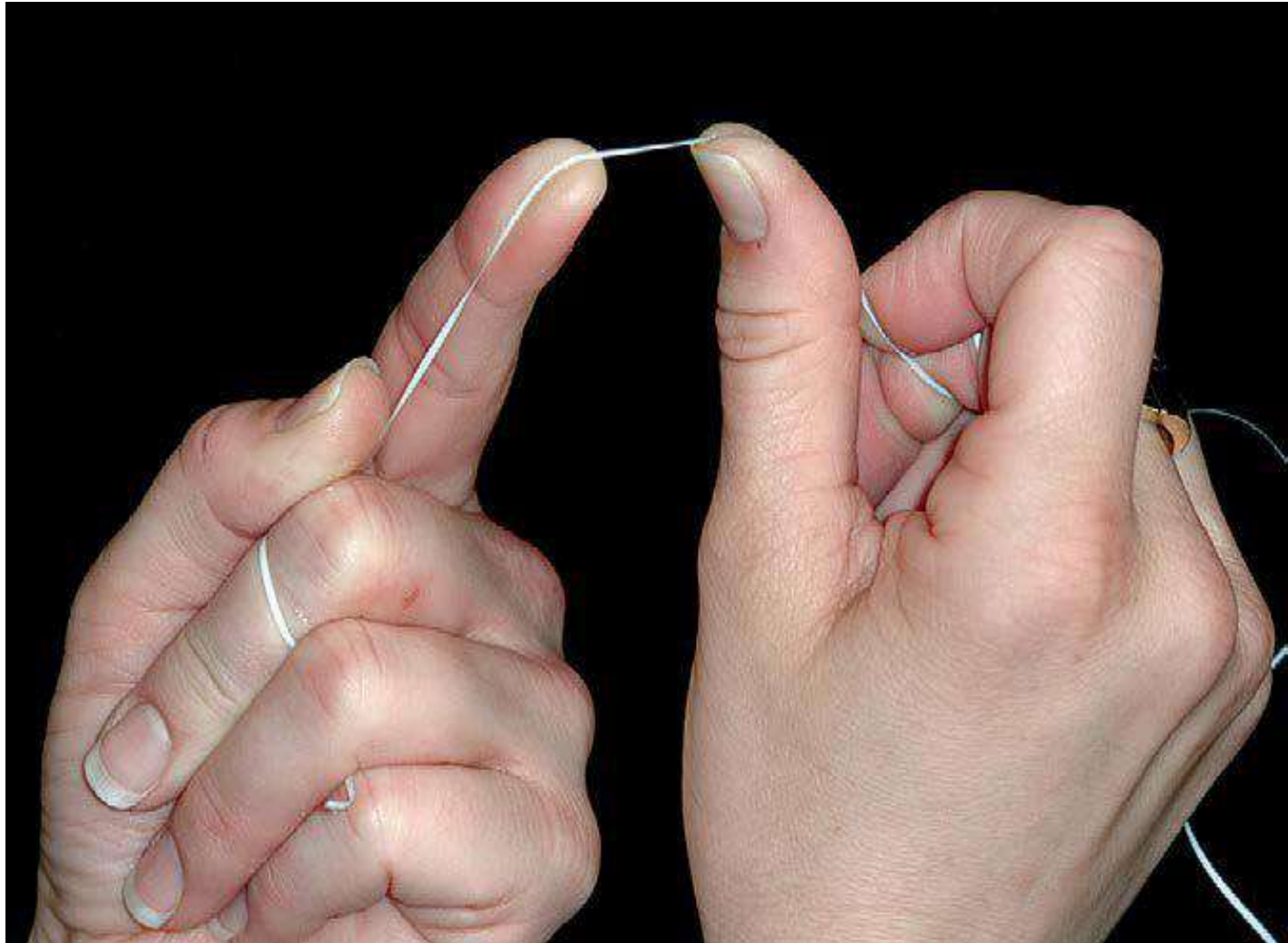
Figure 44-12 Interproximal cleaning devices include wooden tips (A and B), interproximal brushes (C to F), and rubber tip stimulators (G).

1- Dental Floss

Dental floss is the most widely recommended tool for removing plaque from proximal tooth surfaces. Floss is made from nylon filaments or plastic monofilaments, and can be waxed, unwaxed, thick, thin, and even flavored.

Technique. The floss must contact the proximal surface from line angle to line angle to clean effectively. It must also clean the entire proximal surface, including accessible subgingival areas, not just be slipped apical into the contact area.

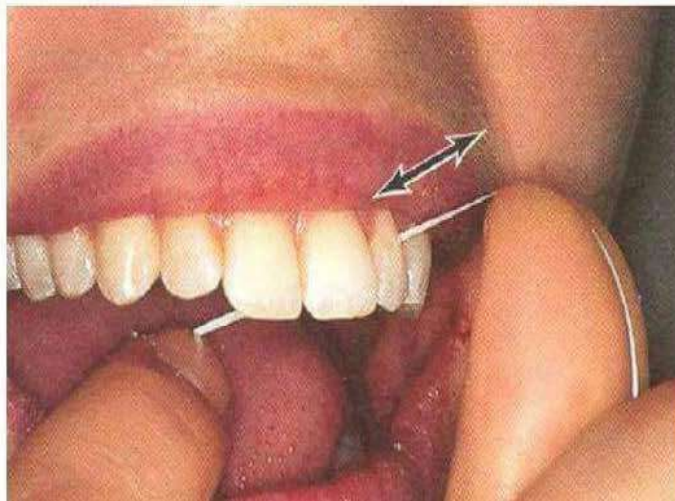
Dental floss should be held securely in the fingers or tied in a loop.



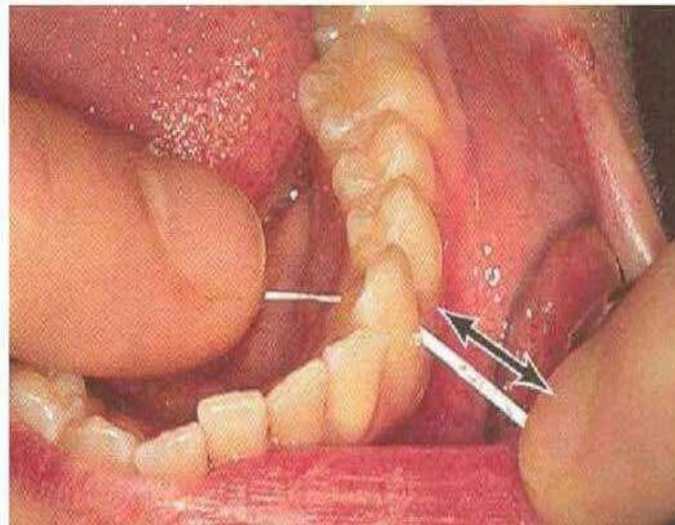
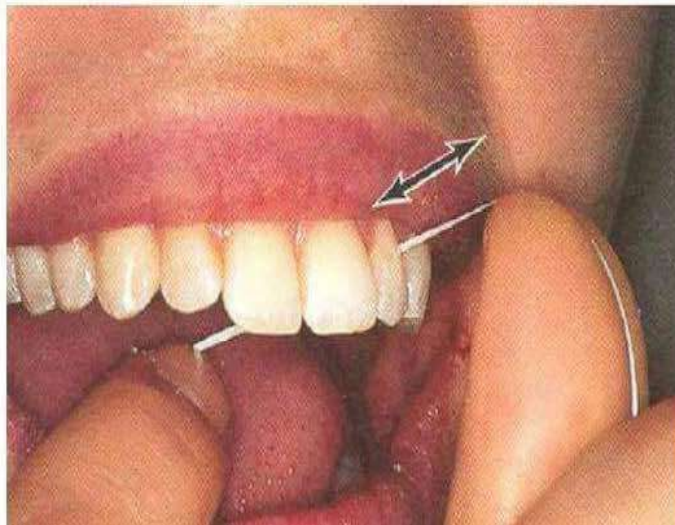
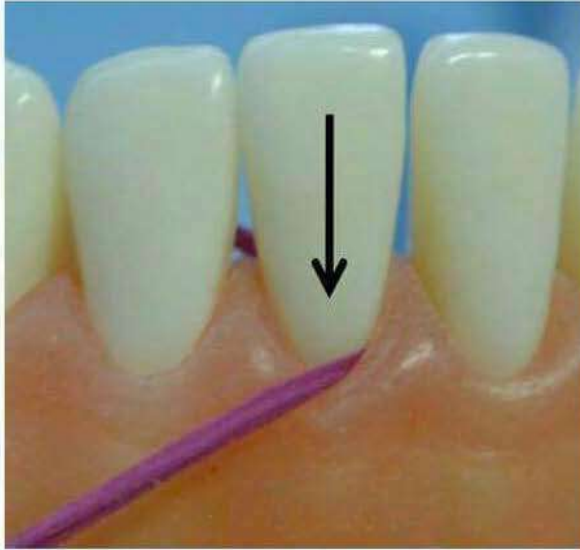
Dental floss should be held securely in the fingers or tied in a loop

1 of 31

Move the floss up and down the tooth



Move the floss up and down the tooth



Dental tape is similar to dental floss only wider



Super floss it is marketed for use in patients with crowns, orthodontics appliances and bridge work. It has a stiffened end to thread between the teeth, a spongy portion to brush the teeth and a floss section for a conventional flossing.





2- Interdental brushes

Concave root surfaces and furcations are often present in periodontal patients who have experienced significant attachment loss and recession, and they are not cleaned well with dental floss. Embrasure spaces vary greatly in size and shape. As a general rule, the larger the space, the larger the device used to clean it should be.

Technique.

Interdental brushes of any style are inserted through interproximal spaces and moved back and forth between the teeth with short strokes.

For the most efficient cleaning, the diameter of brush should be slightly larger than the gingival embrasures to be cleaned. This size permits bristles to exert pressure on both proximal tooth surfaces, working their way into concavities on the roots.

Single-tufted brushes provide access to furcation areas, or isolated areas of deep recession, and work well on the lingual surfaces of mandibular molars and premolars. These areas are often missed when using a toothbrush

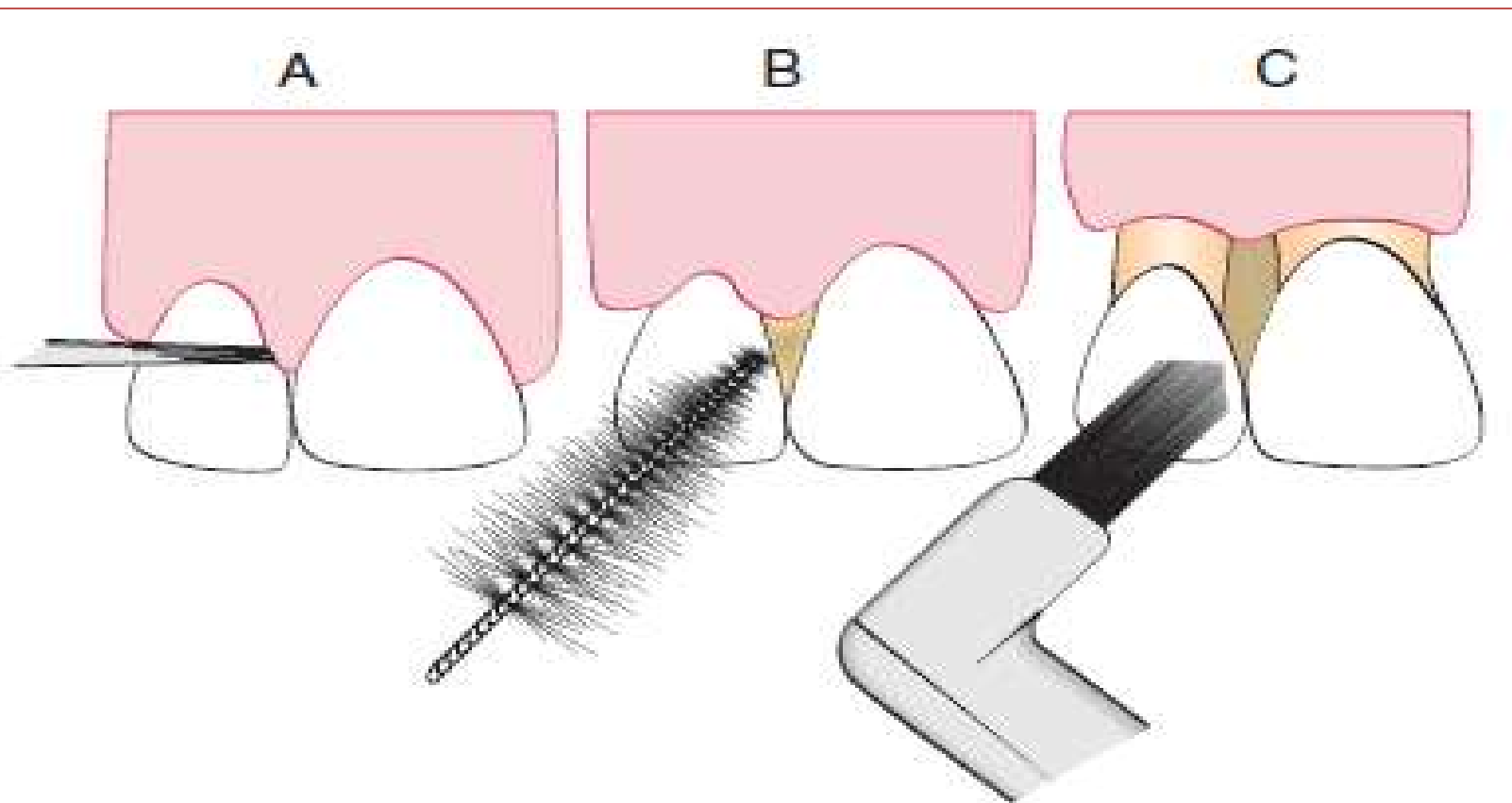


Figure 44-11 Interproximal embrasure spaces vary greatly in patients with periodontal disease. In general, **A**, embrasures with no gingival recession are adequately cleaned using dental floss; **B**, larger spaces with exposed root surfaces require the use of an interproximal brush; and **C**, single-tufted brushes clean efficiently in interproximal spaces with no papillae.

3- Wooden or Rubber Tips

Soft, triangular wooden picks or plastic alternatives are placed in the interdental space with the base of the triangle resting on the gingiva and the sides in contact with the proximal tooth surfaces. The pick is then repeatedly moved in and out of the embrasure to remove plaque. The disadvantage of triangular wooden or plastic tips is that they do not reach well into the posterior areas or on the lingual surfaces and are not shown to remove plaque but can be used to remove food debris.

Wooden toothpick

Triangular wooden tips are also popular with patients. The tip is inserted between the teeth, with the triangular portion resting on the gingival papilla. The tip is moved in and out to remove plaque; however, it is very difficult to use on posterior teeth and from the lingual aspect of all teeth.



Chemical plaque control (Antiseptics):

Adjunctive to mechanical methods; e.g.:
chlorhexidine 0.2% mouth wash.

How should I instruct my patient to use
Chlorhexidinemouth wash?

- 1- Volume used = 10ml.
- 2- Brush your teeth then use the mouth wash after 30 min.
- 3- Rinse your mouth with Chlorhexidine for 60 sec.
- 4- After you spit it, don't eat or drink for at least 30 min. to let the mouth wash takes its action.
- 5- Use it twice daily at morning & evening.
- 6- If you notice any allergic reaction due to its use, stop using it immediately.



Periodontal instruments

**Diagnostic
instrument**



**-dental mirror
-probe
-twizeer**

**Manual
scaling
instrument**



supragingival



**-sickle scaler
-cumine
-push scaler**



subgingival



**Hoe
instrument**

**Root planning
instrument**



**Area specific
curette**

**Universal
cures**

**Polishing
instrument**



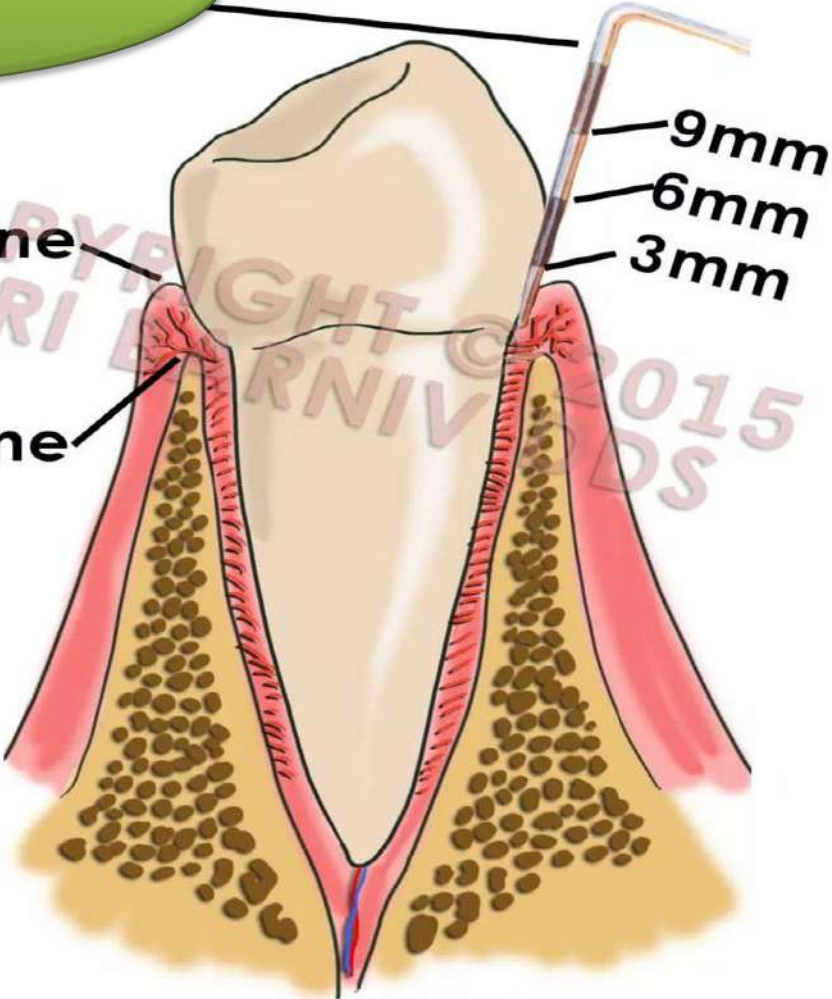
**- rubber cups
- bristles brush
- dental tape**

NORMAL HEALTHY GUMS AND BONE

Periodontal probe

Crest of gum line

Crest of jaw bone



Periodontal probes

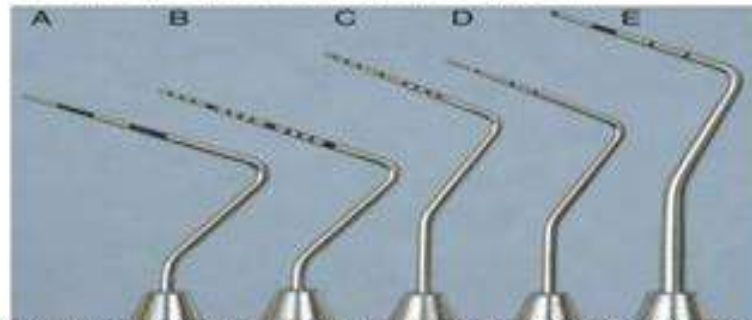
Periodontic Instruments

Explorers

- Use: **1. Locate Sub-gingival Deposits and Caries.**
2. Check Smoothness of root surface after planning.
3. Assess Restorative problems.
- Thin, Flexible, Wire-like working end. Taper to sharp point.
- Curved, Right-angled & Area specific.

Periodontal Probes

- Use: **1. Locate & Measure depth of pocket and determine it's configuration.**
2. Assess Loss of Attachment.
3. Detect Sub-gingival Deposits.
- Tapered with blunt round tip, mm markings for accuracy.
- Ball-end to avoid penetration into junctional epithelium.
- Diameter less than or equal 0.6 mm.
- Probing Force more than 0.25 N traumatize healthy tissue. (25-50g for clinic)
- Ball-end to avoid penetration into junctional epithelium.



A, Marquis color-coded probe. **B**, UHC-15 probe. **C**, University of Michigan "0" probe, with Williams markings. **D**, Michigan "0" probe with markings at 3, 6, and 8 mm. **E**, World Health Organization (WHO) probe.

Periodontal probe for measure depth of sulcus or pocket



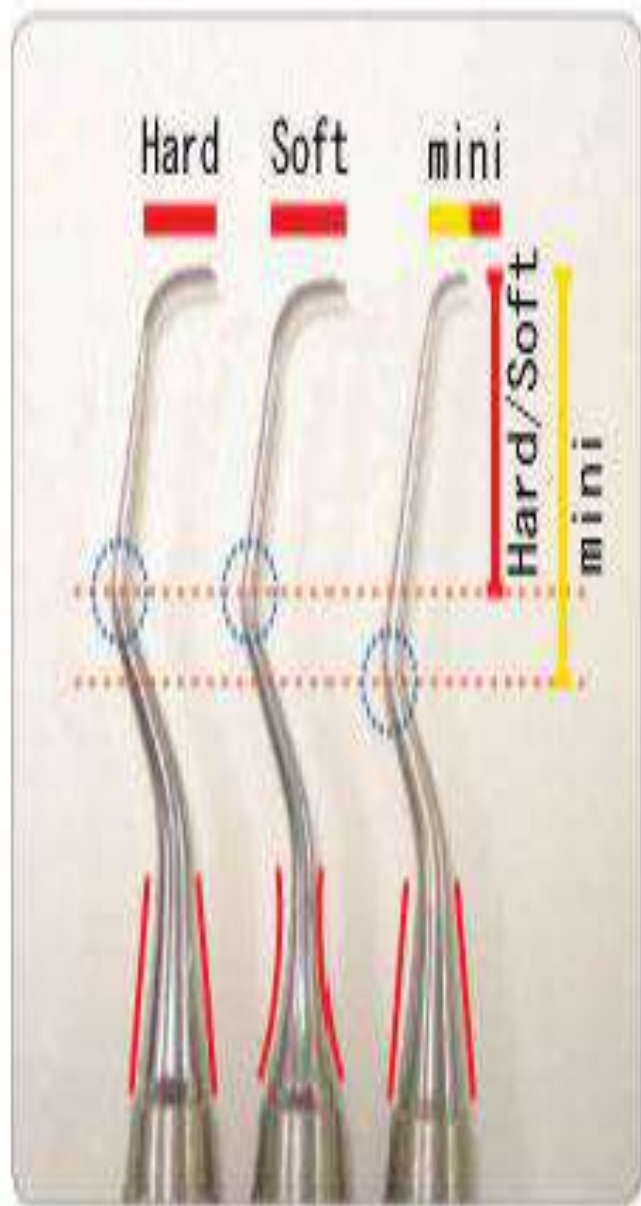
Double Ended Handle #3



Sickle Scaler

Scale scaler for removing of supragingival calculus with pull stroke not used for sugingivally because of caused trauma





curette



Hoe scaler





1.4mm



HSA12-13

• Anterior Hoe Scaler

For anterior buccal and lingual surfaces.

Rubber cups





Brushes for
polishing





Snap-on



Screw type



Latch type

denttp.en.alibaba.com



Tapered



Flat



Cup-shape

Pumice for polishing



Rubber cups for polishing and smoothing



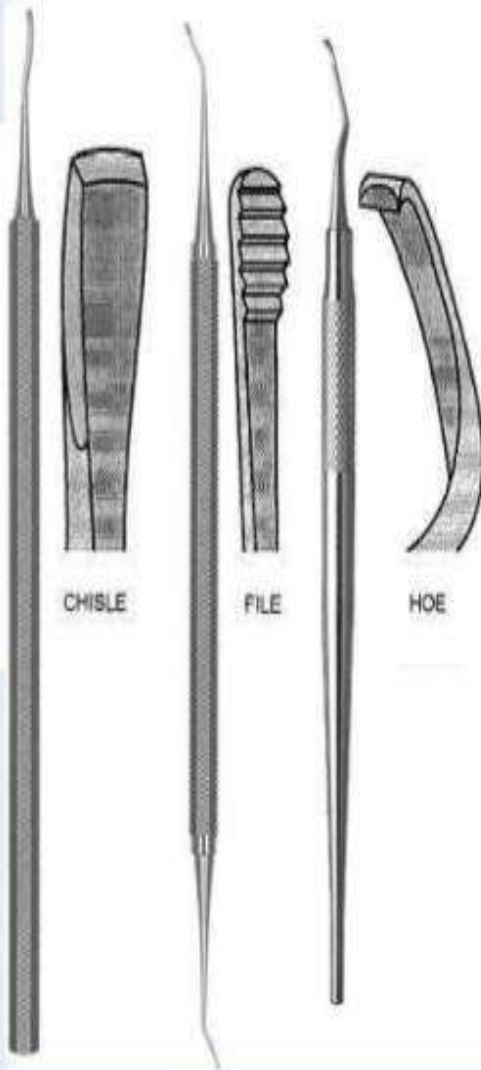


BEFORE



AFTER

SUBGINGIVAL SCALERS



Chisel scaler

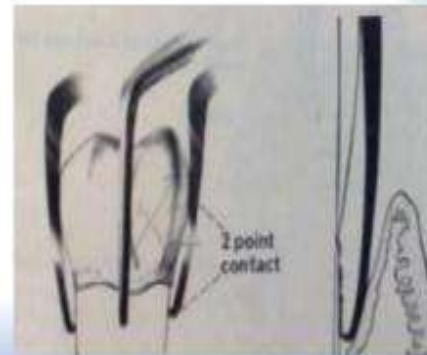
Used in interproximal area
Used in a push motion

File

Used to crush large pieces of
calculus deposits.

Hoe

Efficient in removing subgingival calculus
Blade is beveled at 45 degrees
Working end is bent at an angle of 99
degrees to shank.



Ultrasonic scaler



Ultrasonic scaler



THANKYOU

Dental Calculus

م. سها أسود دهش العزاوي

B.D.S, MSc. Periodontology

- The primary cause of gingival inflammation is bacterial plaque, other predisposing factors including calculus, faulty restoration, orthodontic therapy, use of tobacco, and others.
- **Calculus** consists of mineralized bacterial plaque that forms on the surface of natural teeth and dental prosthesis.

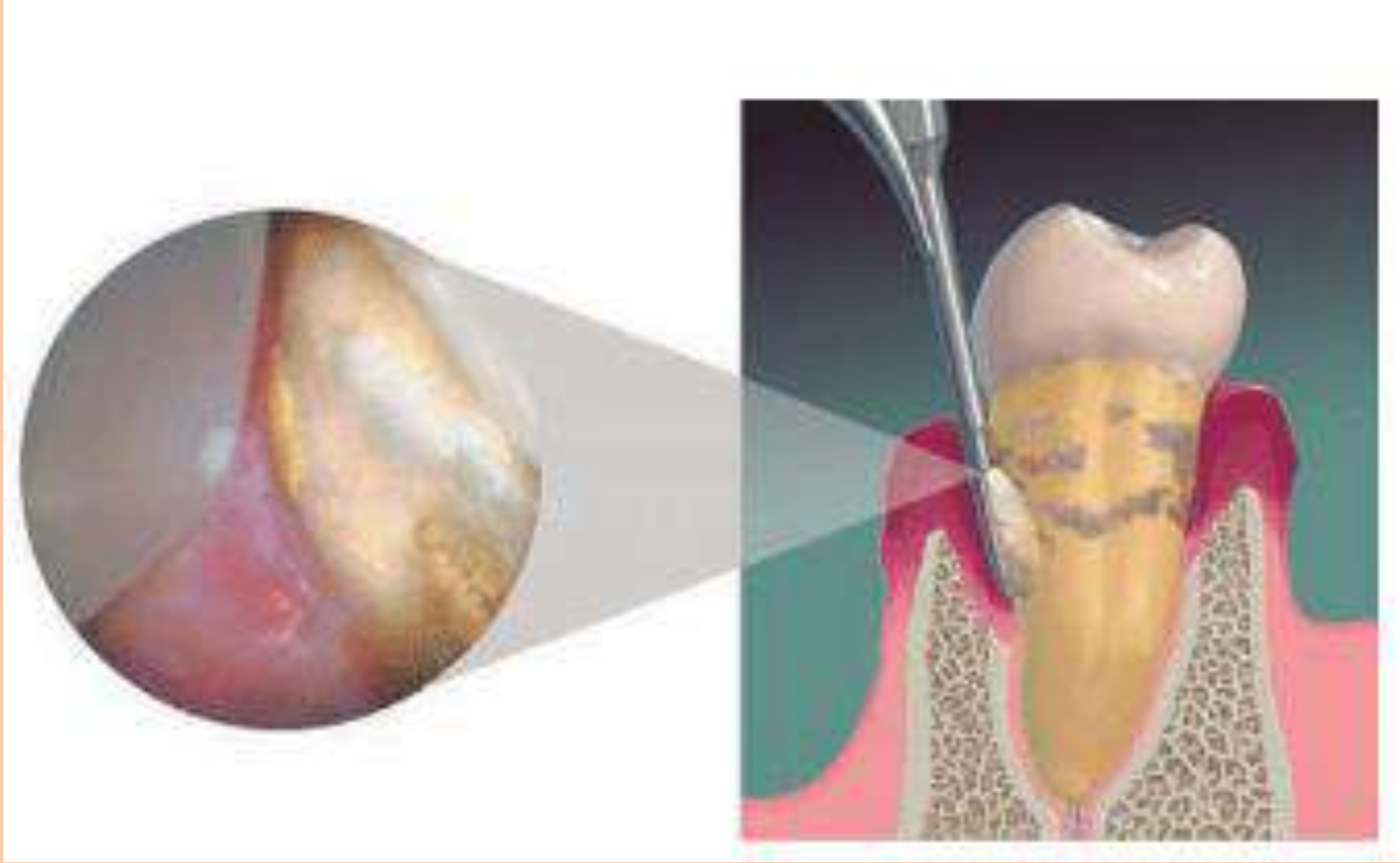
Supra and sub-gingival calculus

- **Supragingival calculus** is located coronal to the gingival margin and therefore visible in the oral cavity,
- usually white to yellowish in colour, hard with claylike consistency and easily detected from the tooth surface, its colour is influenced by contact with such substances as tobacco and food pigments.
- It may localize on a single tooth or group of teeth or it may be generalized throughout the mouth.



- The two most common locations for supragingival calculus to develop are the buccal surface of maxillary molars and the lingual surfaces of mandibular anterior teeth, saliva from the parotid gland flows over the facial surfaces of the upper molars via the parotid duct, while the submandibular and the sublingual glands empty onto lingual surfaces of lower incisors via the submandibular and lingual ducts respectively.

- **Subgingival calculus** is located below the crest of the marginal gingiva and therefore not visible on routine clinical examination,
- the location and extent of the subgingival calculus maybe evaluated by careful tactile perception with a delicate dental instruments such as a dental explorer,
- subgingival calculus is typically hard and dense, frequently appears dark brown or greenish black in colour and firmly attached to the tooth surface.
- When the gingival tissue recede, subgingival calculus becomes exposed and its therefore classified as supragingival.
- Both supra and sub gingival calculus maybe seen by radiograph.



Subgingival calculus detection

Composition

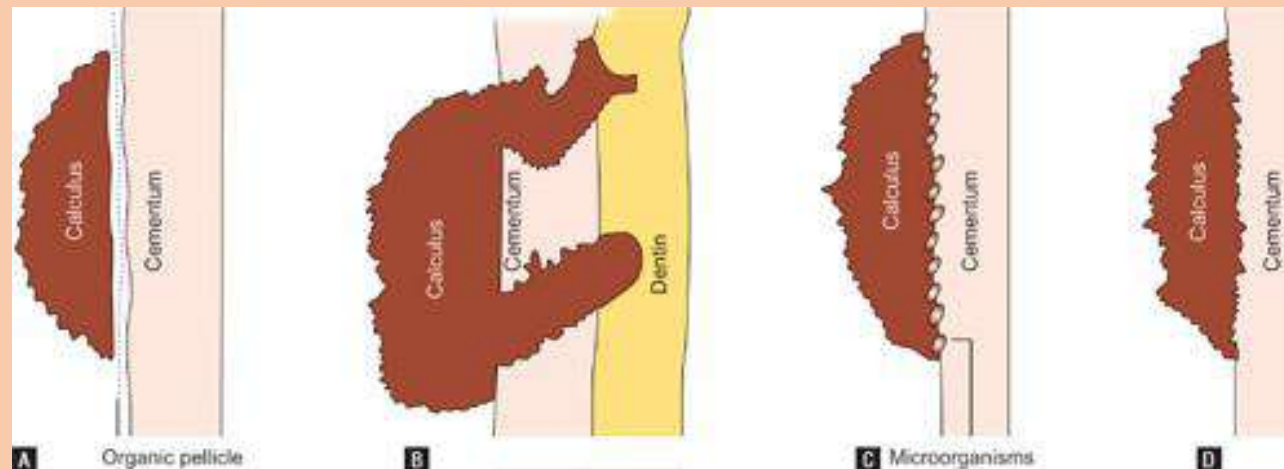
- **Inorganic content** : supragingival calculus consist of inorganic (70% to 90%) and organic components,
- The major inorganic proportions of calculus have been reported as approximately 76% calcium phosphate $\text{Ca}_3(\text{PO}_4)_2$; 3% calcium carbonate CaCO_3 ; traces of magnesium phosphate $\text{Mg}_3(\text{PO}_4)_2$ and other metals .
- At least two thirds of the inorganic component is **crystalline in structure**,
- The four main crystal forms and their approximate percentage are as follows:
- **hydroxyapatite 58%**, **magnesium white-lockite 21%**, octacalcium phosphate 12% and **brushite 9%**.

- Generally two or more crystal forms are typically found in the sample of calculus, **hydroxyapatite and octacalcium phosphate** are detected most frequently in the supragingival calculus and constitute the bulk of the specimen.
- **Brushite** is more common in the mandibular anterior region and
- **magnesium white-lockite** is in the posterior areas,
- The composition of subgingival calculus is similar to that of supragingival calculus with some differences, it has the same hydroxyapatite content, more magnesium white-lockite, and less brushite and octacalcium phosphate

- Organic content: the organic component of calculus consists of a mixture of protein-polysaccharide complexes, desquamated epithelial cells. Leukocytes and various types of microorganisms.
- Salivary proteins present in supragingival calculus are not found subgingivally.

Attachment of calculus to the tooth

- 1. By means of an organic pellicle.
- 2. Mechanical locking into surface irregularities such as carious lesions and resorption lacuna.
- 3. penetration by bacterial calculus into cementum and may appear morphologically similar to cementum and thus termed calculocementum.
- 4. Close adaptation of the undersurface of calculus to depressions or gently sloping surfaces of the unaltered cementum surface.



Formation

- Calculus is dental plaque that has undergone mineralization; the soft plaque is hardened by precipitation of mineral salts which is usually started between first and fourteenth days of plaque formation.
- Calcification started as soon as 4-8 hours, calcifying plaque may become 50% mineralized in 2 days, 60%-90% mineralized in 12 days. However, the formation of dental calculus with the mature crystalline composition of old calculus may require months to years.
- Saliva is the source of mineralization for supra gingival calculus whereas the serum transudate (Gingival crevicular fluid) is source of mineralization of sub gingival calculus.
- Microorganisms are not always essential in calculus formation because calculus occurs readily in germ free rodents,

- The initiation of calcification and rate of accumulation vary among teeth in same individual, so person may be heavy, moderate or slight calculus former.
- Calculus formation continues until it reaches maximum after which it reduced in amount due to mechanical wear from food and the cheeks, lip and tongue, also the use of anti-calculus (anti tarter) dentifrices reduce both quality and quantity of calculus.

Theories regarding the mineralization of Calculus

- The theoretical mechanisms by which plaque becomes mineralized can be stratified into two categories:

1. Mineral precipitation results from a local rise in the degree of saturation of calcium and phosphate ions, which may be brought about in the following several ways:

- A rise in the pH of the saliva causes precipitation of calcium phosphate salts by lowering the precipitation constant. The pH may be elevated by the loss of carbon dioxide and the formation of ammonia by dental plaque bacteria or by protein degradation.

- Colloidal proteins in saliva bind calcium and phosphate ions and maintain a supersaturated solution with respect to calcium phosphate salts.
- Phosphatase liberated from dental plaque, desquamated epithelial cells, or bacteria precipitates calcium phosphate by hydrolyzing organic phosphates in saliva, thus increasing the concentration of free phosphate ions.

- **2. Seeding agents induce small foci of calcification that enlarge and coalesce to form a calcified mass.** This concept has been referred to as the **epitactic concept** or more appropriately, hetero- geneous nucleation. The seeding agents in calculus formation are not known, but it is suspected that the intercellular matrix of plaque plays an active role. The carbohydrate-protein complexes may initiate calcification by removing calcium from the saliva (chelation) and binding with it to form nuclei that induce subsequent deposition of minerals.

Etiologic Significance


- The non-mineralized plaque on the calculus surface is the principle irritant for initiating gingivitis.
- The underlying calcified portion is a significant contributing factor since it provides a fixed nidus for the continued accumulation of plaque and remains it close to gingiva.
- calculus plays an important role in maintaining periodontal diseases by keeping plaque in close contact with the gingival tissue and creating area where plaque removal is impossible unless we remove calculus.
- it is a secondary etiologic factor for periodontitis and it is the most prominent plaque retentive factor which has to be removed as a basis for adequate periodontal therapy and prophylactic activities..Removal of supra and sub gingival plaque and calculus constitute the cornerstone of periodontal therapy.

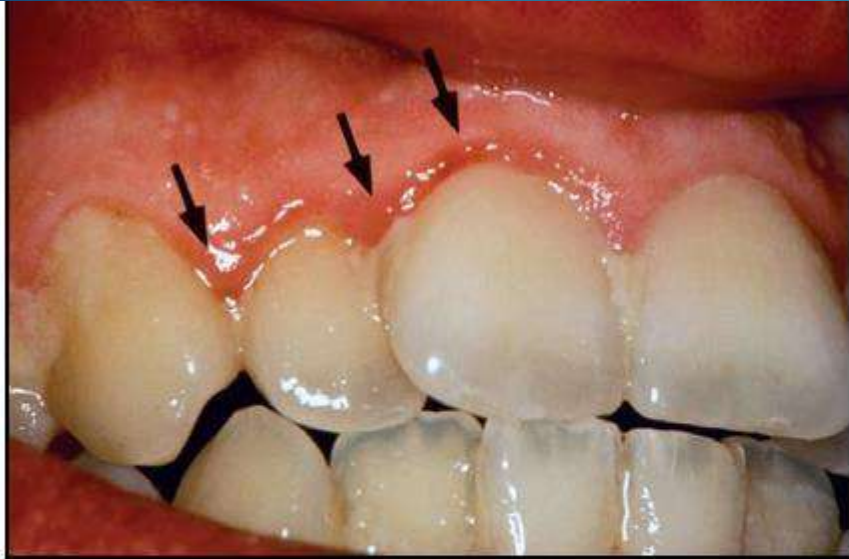
Thank you

DENTAL PLAQUE BIOFILM

م. سها أسود دهش العزاوي

B.D.S, MSC. PERIODONTOLOGY

- 
- ▶ Oral biofilms are **functionally and structurally** organized polymicrobial communities that are embedded in an extracellular matrix of exopolymers on mucosal and dental surfaces.
 - ▶ Dental plaque is defined **clinically** as a structured resilient yellow-grayish biofilm that adheres firmly to the intraoral hard surfaces, including removable and fixed restorations. The tough extracellular matrix makes it impossible to remove plaque by rinsing or the use of sprays. Plaque can thus be differentiated from other deposits that may be found on the tooth surface such as materia alba and calculus.
 - ▶ **Microorganisms represent the main component of dental plaque, with approximately 10^{11} bacteria** contained in one gram of plaque (wet weight)



Clinical picture of 10-day-old supragingival plaque. Black rows indicate early signs of gingival inflammation.



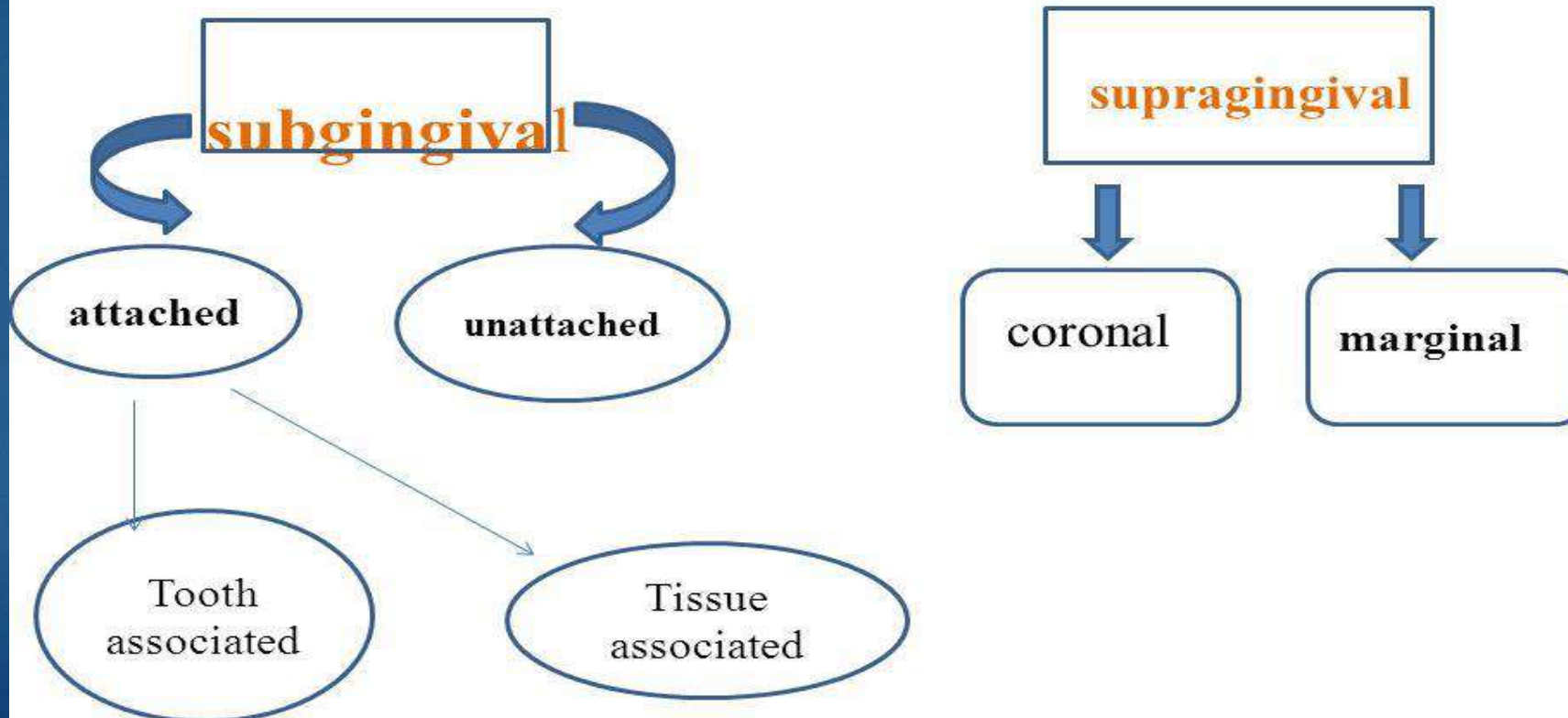
Supragingival calculus is depicted on the buccal surface of maxillary molars adjacent to the orifice for the parotid duct.

	Materia alba	Dental plaque	Claculus
1	White cheese-like accumulations	Resilient clear to yellow grayish	Hard deposits formed by mineralization of dental plaque
2	A soft accumulation of salivary proteins	Primarily composed of bacteria in a matrix of salivary proteins	
3	Lack of organized structure (not complex) as dental plaque)	Considered as a biofilm	Generally covered by a layer of un-mineralized dental plaque
4	Easily displaced by a water spray	Removed only by mechanical rinsing (tooth brushing)	

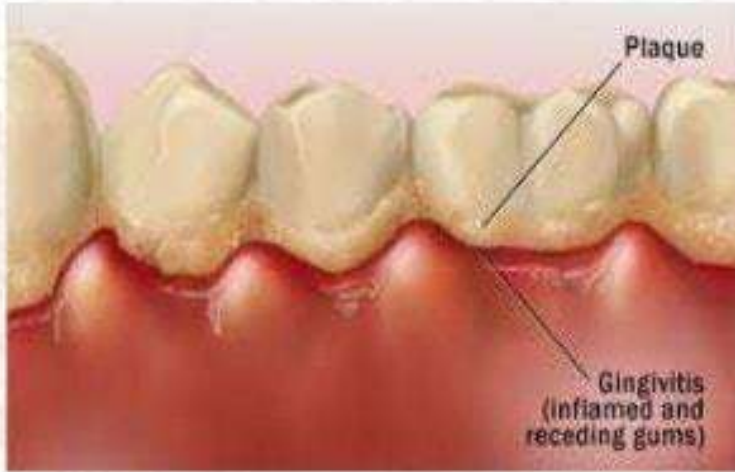
Classification of Plaque

Classification of plaque:

- According to tooth surface: (location)



SUPRAGINGIVAL PLAQUE



Tooth-Attached Plaque

Mostly gram+ bacteria; with some gram- cocci and rods.

Tooth attached plaque

Unattached plaque

Epithelial associated plaque

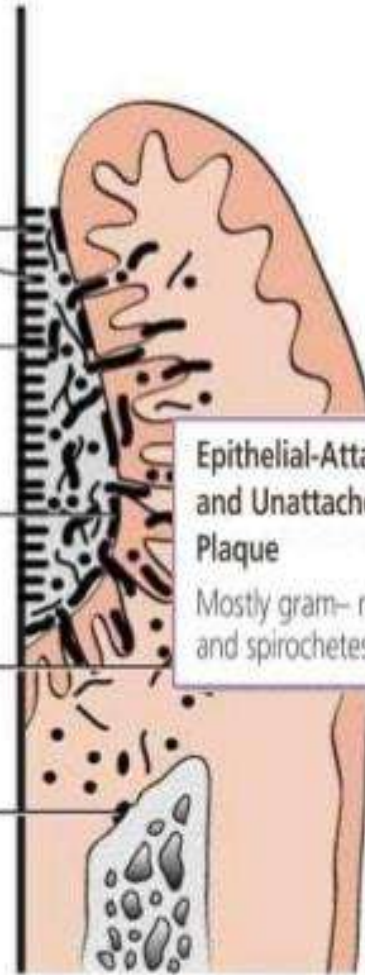
Bacteria within connective tissue


Bacteria on bone surface

SUBGINGIVAL PLAQUE

Epithelial-Attached and Unattached Plaque

Mostly gram- rods and spirochetes.



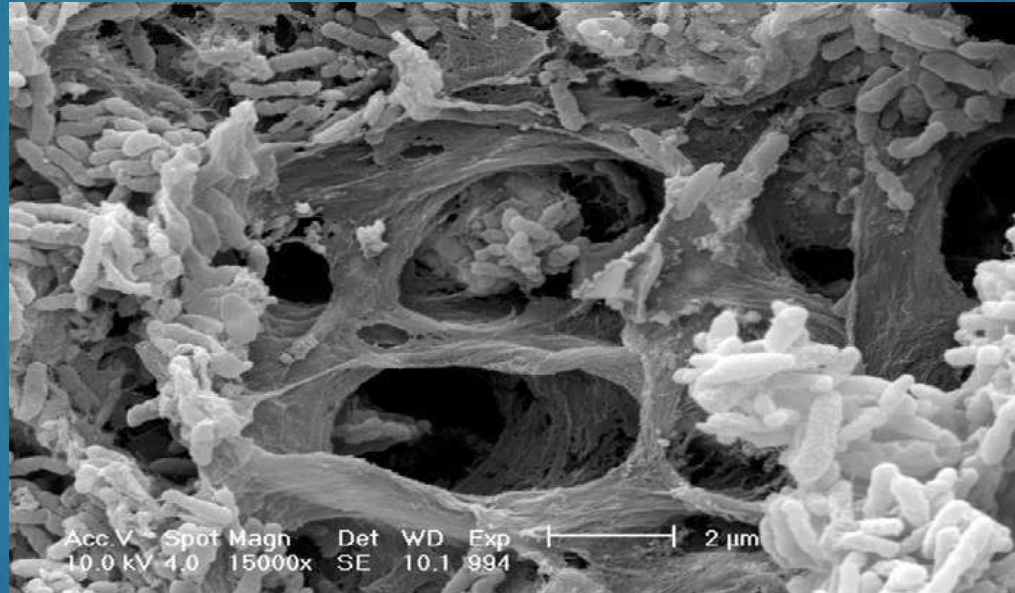


Supragingival plaque typically demonstrates a **stratified organization** of a multilayered accumulation of bacterial morphotypes. **Gram-positive** cocci and short rods predominate at the tooth surface, whereas **gram-negative** rods and filaments, as well as spirochetes, predominate in the outer surface of the mature plaque mass


In general, the subgingival microbiota differs in composition from the supragingival plaque, primarily because of the **local availability of blood products** and a **low reduction-oxidation (redox) potential**, which characterizes the **anaerobic** environment.

- ▶ Both morphologic and microbiologic studies of subgingival plaque reveal distinctions between the tooth-associated and soft tissue-associated regions of subgingival plaque.
- ▶ The tooth-associated cervical plaque, adhering to the root cementum, does not markedly differ from that observed in gingivitis. At this location, filamentous microorganisms dominate, but cocci and rods also occur. This plaque is dominated by gram positive rods and cocci, including *S. mitis*, *S. sanguinis*, *Actinomyces oris*. However, in the deeper parts of the pocket, the filamentous organisms become fewer in numbers, and in the apical portion they seem to be virtually absent. Instead, the microbiota is dominated by smaller organisms
- ▶ The apical border of the plaque mass is separated from the junctional epithelium by a layer of host leukocytes

- ▶ The layers of microorganisms facing the soft tissue lack a definite intermicrobial matrix and contain primarily gram-negative rods and cocci as well as large numbers of filaments, flagellated rods, and spirochetes. Host tissue cells (e.g., white blood cells and epithelial cells) may also be found in this region. Bacteria are also found within the host tissues, such as in the soft tissues and within epithelial cells, as well as in the dentinal tubules.

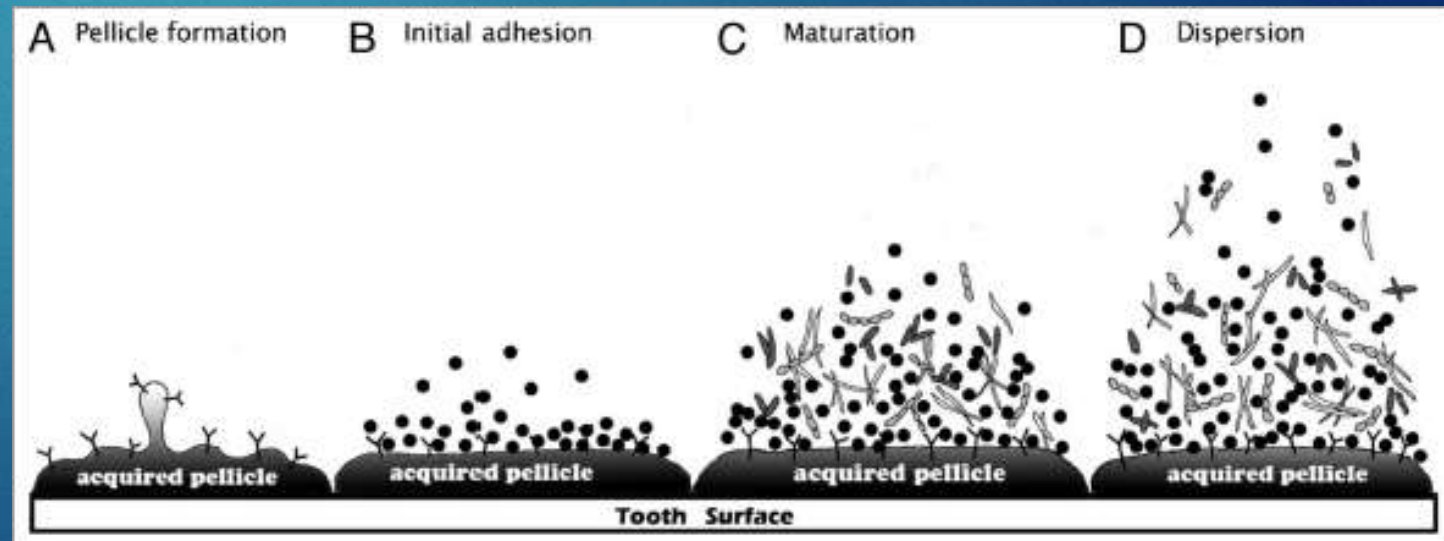


- The composition of the subgingival plaque depends on pocket depth; the apical part is more dominated by spirochetes, cocci and rods, whereas in the coronal part more filaments are observed.

- 
- ▶ The site specificity of plaque is significantly associated with diseases of the periodontium.
 - ▶ **Marginal plaque**, for example, is of prime importance in the initiation and development of **gingivitis**.
 - ▶ **Supragingival plaque and tooth-associated subgingival plaque** are critical in **calculus** formation and **root caries**; whereas
 - ▶ **Tissue associated subgingival plaque** is important in the tissue destruction that characterizes different forms of **periodontitis**.

Accumulation of a Dental Plaque Biofilm

- ▶ The process of plaque formation can be divided into several phases:
- ▶ 1- The formation of the pellicle on the tooth surface.
- ▶ 2- Initial adhesion/attachment of bacteria.
- ▶ 3- Colonization/plaque maturation.



Formation of the Pellicle

- ▶ It is translucent, homogenous, thin film covering all surfaces in the oral cavity, formed by adsorption of proteins on tooth surface.
- ▶ The pellicle on tooth surface consists of more than 180 peptides, proteins, and glycoproteins, and other molecules that can function as adhesion sites (receptors) for bacteria.
- ▶ Salivary pellicle can be detected on clean enamel surfaces within 1 minute.
- ▶ Bacteria that adhere to tooth surfaces do not contact the enamel directly but interact with the acquired enamel pellicle.

SIGNIFICANCE OF PELLICLE:

- **PROTECTIVE:** provide barrier against acids thus may reduce dental caries attack.
- **LUBRICATION:** keep surface moist prevent drying.
- **NIDUS FOR BACTERIA:** Plaque formation by adherence of microorganisms.
- **ATTACHEMENT OF CALCULUS:** A mode of calculus attachment in acquired pellicle.

Initial Adhesion/Attachment of Bacteria.

- ▶ The initial steps of transport and interaction with the surface are essentially nonspecific (i.e., they are the same for all bacteria).
- ▶ Specific interactions between microbial cell surface “adhesin” molecules and receptors in the salivary pellicle that determines whether a bacterial cell will remain associated with the surface.
- ▶ small proportion of oral bacteria possess adhesins that interact with receptors in the host pellicle, and these organisms are generally the most abundant bacteria in biofilms on tooth enamel shortly after cleaning. These species are considered the “primary colonizers” of tooth surfaces.
- ▶ The primary colonizers provide new binding sites for adhesion by other oral bacteria.

Colonization and Plaque Maturation.

- ▶ The primary colonizing bacteria adhered to the tooth surface provide a new receptors for attachment by other bacteria in a process known as “coadhesion.”
- ▶ Together with growth of adherent microorganisms, coadhesion leads to the development of micro colonies and eventually to a mature biofilm.
- ▶ Different species or even different strains of a single species have distinct sets of coaggregation partners. Fusobacteria coaggregate with all other human oral bacteria while Veillonella spp., Capnocytophaga spp. and Prevotella spp. bind to streptococci and/or actinomyces. Each newly cell becomes itself a new surface and therefore, may act as a coaggregation bridge to the next potentially accreting cell type that passes by.

- ▶ Secondary colonizers ,such as *Prevotella intermedia*, *Prevotella loescheii*, *Capnocytophaga spp.*, *F. nucleatum*, and *P. gingivalis* do not initially colonize clean tooth surfaces but adhere to bacteria already in the plaque mass.

COMPOSITION OF PLAQUE:

COMPOSED mainly of:

- Microorganisms ,
- Intermicrobial matrix

(a) Bacterial portion:

- 70 to 80 % of total solid plaque volume.
- 1 g (WET WT) contains approx. 10^{11} bacteria.

(B) Intermicrobial Matrix

- the material present between the bacteria in the dental plaque is called the intermicrobial matrix.
- accounts for approx 25% of plaque volume.

Three sources may contribute to the intermicrobial matrix:

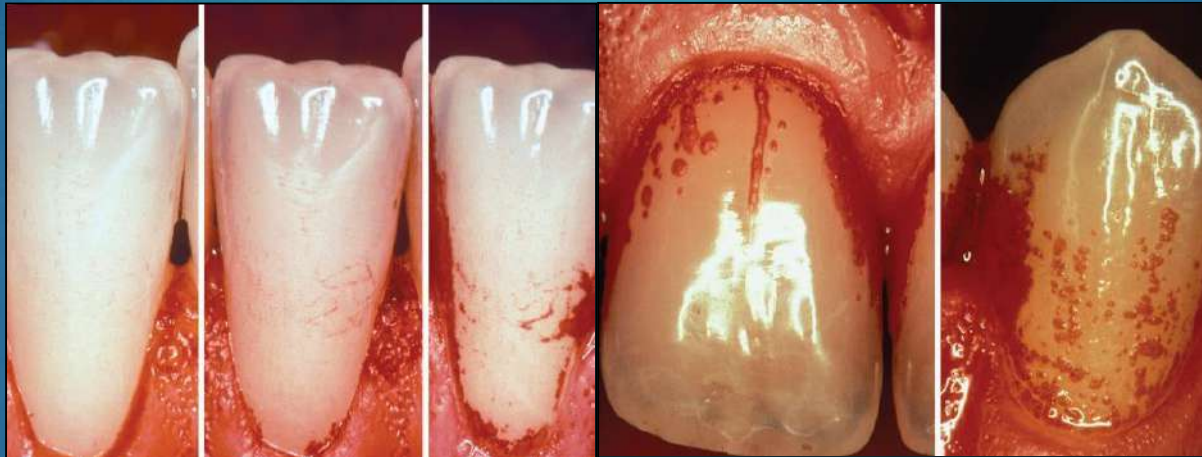
- The plaque microorganisms
- The saliva
- Gingival exudates

Factors Affecting Supragingival Dental plaque formation

- ▶ During the **first 24 hours** starting from a clean tooth surface, plaque growth is nearly undetectable clinically.
- ▶ **after 4 days**, on average **30%** of the total coronal tooth area will be covered with plaque.
- ▶ the microbial composition of the dental plaque will change with a **shift from the early aerobic environment** characterized by **gram-positive facultative species** to a **highly oxygen-deprived environment** in which **gram negative anaerobic microorganisms** predominate.

Topography of Supragingival Plaque

- ▶ initial growth along the gingival margin and from the interdental space. Later, a further extension in the coronal direction.
- ▶ Plaque formation can also start from grooves, cracks, or pits.



Irregular plaque growth patterns follow tooth surface irregularities

Surface Micro roughness.

- ▶ Rough intraoral surfaces (e.g. crown margins, implant abutments, and denture bases) accumulate and retain more plaque and calculus in terms of thickness, area, and colony-forming units. Ample plaque also reveals an increased maturity/pathogenicity of its bacterial components, characterized by an increased proportion of motile organisms and spirochetes and/or a denser packing of them. Smoothing an intraoral surface decreases the rate of plaque formation.

Individual Variables Influencing Plaque Formation.

- ▶ The rate of plaque formation differs significantly between subjects,
- ▶ A distinction is often made between “heavy” (fast) and “light” (slow) plaque formers.

Variation within the Dentition.

Early plaque formation occurs faster:

- ▶ in the lower jaw (when compared to the upper jaw);
- ▶ in molar areas; on the buccal tooth surfaces when compared to palatal sites (especially in the upper jaw); and
- ▶ in the interdental regions when compared to the buccal or lingual surfaces.

Impact of Gingival Inflammation and Saliva

- ▶ Several studies clearly indicate that **plaque formation is more rapid on tooth surfaces facing inflamed gingival margins** than on those adjacent to healthy gingival. These studies suggest that the increase in crevicular fluid production enhances plaque formation. Probably, some substance(s) from this exudate (e.g. minerals, proteins, or carbohydrates) favor both the initial adhesion and/or the growth of the early colonizing bacteria.
- ▶ Additionally, it is known that during the night, plaque growth rate is reduced by some 50%. This seems surprising, since one would expect that reduced plaque removal and the decreased salivary flow at night would enhance plaque growth. The fact that the supragingival plaque obtains its nutrients mainly from the saliva appears to be of greater significance than the antibacterial activity of saliva.

The Impact of Patient's Age.

- ▶ Although older studies were contradictory, more recent papers clearly indicate that a subject's age does not influence de novo plaque formation.

Spontaneous Tooth Cleaning.

- ▶ Many clinicians still believe that plaque is removed spontaneously from the teeth such as during eating. However, based on the firm attachment between bacteria and surface, this seems unlikely. Even in the occlusal surfaces of the molars, plaque remains, even after chewing fibrous food.



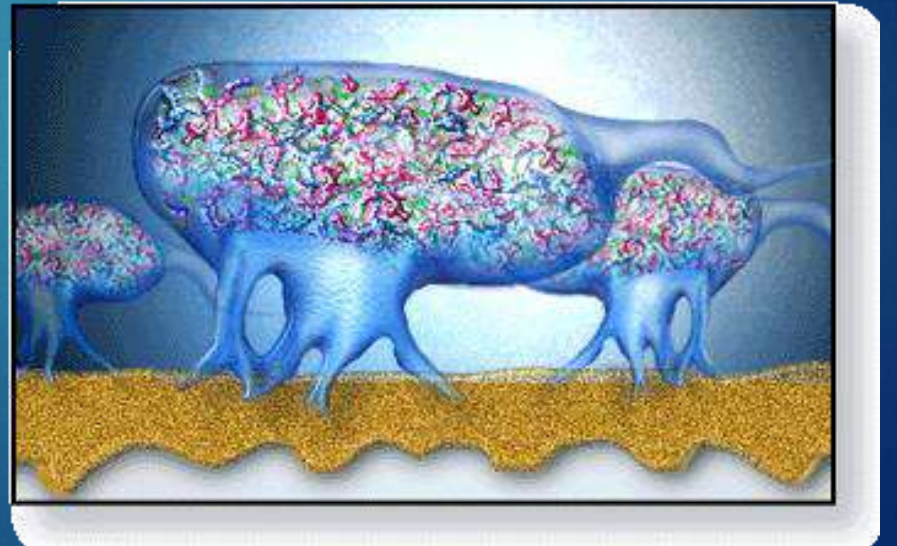
No reduction of 100 hours old dental plaque before dinner (**A**), and after dinner (**B**) with eating fibrous food

Metabolism of Dental Plaque Bacteria

- ▶ The majority of nutrients for dental plaque bacteria originate from saliva or GCF.
- ▶ Although the host diet provides an occasional but nevertheless important food supply.
- ▶ The growth of *P. gingivalis* is enhanced by metabolic byproducts produced by other microorganisms, such as succinate from *Capnocytophaga ochracea* and protoheme from *Campylobacter rectus*. Overall, the total plaque population is more efficient than any one constituent organism at releasing energy from the available substrates.
- ▶ Metabolic interactions occur also between the host and plaque microorganisms. Increases in steroid hormones are associated with significant increases in the proportions of *P. intermedia* found in subgingival plaque. These nutritional interdependencies are probably critical to the growth and survival of microorganisms in dental plaque and may partly explain the evolution of highly specific structural interactions observed among bacteria in plaque.

PLAQUE AS A BIOFILM

- As the bacteria attach to a surface and to each other, they cluster together to form microcolonies that are attached to the surface.
- Each microcolony is a tiny, independent community containing thousands of compatible bacteria.
- Different microcolonies may contain different combinations of bacterial species.



Communication between Biofilm Bacteria

- ▶ Bacterial cells do not exist in isolation. In a biofilm, bacteria have the capacity to communicate with each other. One example of this is **quorum sensing**, in which bacteria secrete a signaling molecule that accumulates in the local environment and triggers a response such as a change in the expression of specific genes once they reach a critical threshold concentration. The **threshold concentration** is reached only at a high-cell density, and therefore bacteria sense that the population has reached a critical mass, or quorum. Responses are induced only when a threshold concentration of the peptide is attained, and thus the peptides act as cell density, or quorum, sensors.

Biofilms and Antimicrobial Resistance

- ▶ organisms in a biofilm are 1000 to 1500 times more resistant to antibiotics than in their planktonic state.
- ▶ The mechanisms of this increased resistance differ from species to species, from antibiotic to antibiotic, and for biofilm growing in different habitats.
- ▶ An important mechanism of resistance appears to be the **slower rate of growth** of bacterial species in a biofilm, which makes them less susceptible to many but not all antibiotics. The **biofilm matrix**, although not a significant physical barrier to the diffusion of antibiotics, does have certain properties that can retard antibiotic penetration
- ▶ In addition, **extracellular enzymes** such as β -lactamases, formaldehyde lyase, and formaldehyde dehydrogenase may become trapped and concentrated in the extracellular matrix, thus inactivating some antibiotics

Thank you



Dental Stain

م. سها أسود دهش العزاوي

B.D.S, MSc. Periodontology

Dental Stain

- Pigmented deposits on the tooth surface.
- There has been a recent increase in interest in the treatment of tooth staining and discolorations as shown by the large number of tooth whitening agents appearing on the market.
- The correct diagnosis for the cause of discoloration is important as, invariably, it has a profound effect on treatment outcomes.

COLOUR AND COLOUR PERCEPTION

- Teeth are typically composed of a number of colours and a gradation of colour occurs in an individual tooth from the gingival margin to the incisal edge of the tooth. The gingival margin often has a darker appearance because of the close approximation of the dentine below the enamel.
- In most people canine teeth are darker than central and lateral incisors and younger people characteristically have lighter teeth, particularly in the primary dentition. Teeth become darker as a physiological age change; this may be partly caused by the **laying down of secondary dentine**, incorporation of **extrinsic stains** and **gradual wear of enamel** allowing a greater influence on colour of the underlying dentine.

CLASSIFICATION OF TOOTH DISCOLOURATION

- **INTRINSIC DISCOLORATION**

- Intrinsic discoloration occurs following a change to the structural composition or thickness of the dental hard tissues.
- A number of metabolic diseases and systemic factors are known to affect the developing dentition and cause discoloration as a consequence. Local factors such as injury are also recognized:
 - 1. Alkaptonuria
 - 2. Congenital erythropoietic porphyria
 - 3. Congenital hyperbilirubinaemia
 - 4. Amelogenesis imperfecta
 - 5. Dentinogenesis imperfecta

6. Tetracycline staining
7. Fluorosis
8. Enamel hypoplasia
9. Pulpal haemorrhagic products
10. Root resorption
11. Ageing

- **EXTRINSIC DISCOLORATION**

- Extrinsic discoloration is outside the tooth substance and lies on the tooth surface or in the acquired pellicle. The origin of the stain may be:
 - 1. Metallic
 - 2. Non-metallic

- **INTERNALIZED DISCOLORATION**

- Internalized discoloration is the incorporation of extrinsic stain within the tooth substance following dental development. It occurs in enamel defects and in the porous surface of exposed dentine. The routes by which pigments may become internalized are:

1. Developmental defects

2. Acquired defects

- a) Tooth wear and gingival recession

- b) Dental caries

- c) Restorative materials

THE MECHANISMS OF TOOTH DISCOLORATION

- **INTRINSIC TOOTH DISCOLORATION** occurs during tooth development and results in an alteration of the light transmitting properties of the tooth structure.
- *1. Alkaptonuria:* This inborn error of metabolism results in incomplete metabolism of tyrosine and phenylalanine, which promotes the buildup of homogentisic acid. This affects the permanent dentition by causing a **brown discolouration**.

- *2. Congenital erythropoietic porphyria:*
- Metabolic disorder in which there is an error in porphyrin metabolism leading to the accumulation of porphyrins in bone marrow, red blood cells, urine, faeces and teeth causes A **red-brown discolouration** of the teeth.
- *3. Congenital hyperbilirubinaemia:* The breakdown products of haemolysis will cause a **yellow-green** discolouration.
- *4. Amelogenesis imperfecta:* In this hereditary condition, enamel formation is disturbed with regard to mineralization or matrix formation and is classified accordingly. The appearance depends upon the type of amelogenesis imperfecta, varying from the relatively mild hypomature 'snow-capped' enamel to the more severe hereditary hypoplasia with thin, hard enamel which has a **yellow to yellow-brown appearance**.



Amelogenesis Imperfecta

- *5. Dentinogenesis imperfecta*: Dentine defects may occur genetically or through environmental influences.
- The teeth are usually **bluish or brown** in color, and demonstrate opalescence on transillumination. The pulp chambers often become obliterated and the dentine undergoes rapid wear, once the enamel has chipped away, to expose the amelo-dentinal junction.



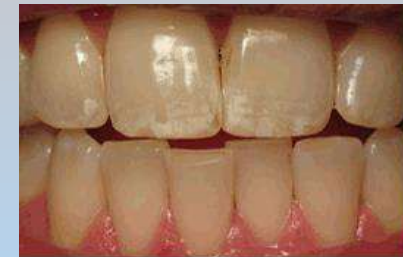
Figure 1. *Generalized opalescent tooth discoloration.*

- *6. Tetracycline staining:* Systemic administration of tetracyclines during development is associated with deposition of tetracycline within bone and the dental hard tissues. Tetracycline is able to cross the placental barrier and should be avoided from 29 weeks *in utero* until full term to prevent incorporation into the dental tissues. Since the permanent teeth continue to develop in the infant and young child until 12 years of age, tetracycline administration should be avoided in children below this age and in breast-feeding and expectant mothers. Teeth affected by tetracycline have a **yellowish** or **brown-grey** appearance



- *7. Fluorosis:*

- This may arise endemically from naturally occurring water supplies or from fluoride delivered in mouth rinses, tablets or toothpastes as a supplement. The severity is related to age and dose.
- The enamel is often affected and may vary from areas of flecking to diffuse opacous mottling, whilst the color of the enamel ranges from chalky white to a dark brown/black appearance. The brown/black discoloration is posteruptive and probably caused by the internalization of extrinsic stain into the porous enamel. Fluoride only causes fluorosis in concentrations of greater than 1 ppm in drinking water.



“Very Mild”



“Mild”



“Moderate”



“Severe”

- *8. Enamel hypoplasia:* This condition may be localized or generalized. The most common localized cause of enamel hypoplasia is likely to occur following **trauma or infection in the primary dentition**. Such localized damage to the tooth-germ will often produce a hypoplastic enamel defect, which can be related chronologically to the injury. Disturbance of the developing tooth germ may occur in a large number of fetal or maternal conditions eg maternal vitamin D deficiencies, rubella infection, drug intake during pregnancy and in pediatric hypocalcaemic conditions. Such defects will be chronologically laid down in the teeth depending on the state of development at the time of interference. There may be pitting or grooving which predisposes to extrinsic staining of the enamel.

- *9. Pulpal hemorrhagic products:*
- The discoloration of teeth following severe trauma was considered to be caused by pulpal hemorrhage.
- Haemolysis of the red blood cells would follow and release the hem group to combine with the putrefying pulpal tissue to form black iron sulphide.
- The depth of dentinal penetration determines the degree of discolouration.
- *10. Ageing:* The natural laying down of secondary dentine affects the light-transmitting properties of teeth resulting in a gradual darkening of teeth with age.

EXTRINSIC DISCOLORATION

- The causes of extrinsic staining can be divided into two categories; those compounds which are incorporated into the pellicle and produce a stain as a result of their basic colour, and those which lead to staining caused by chemical interaction at the tooth surface.
- Direct staining has a multi-factorial aetiology with chromogens derived from dietary sources or habitually placed in the mouth. Tobacco smoking and chewing are known to cause staining, as are particular beverages such as tea and coffee.
- Indirect extrinsic tooth staining is associated with cationic antiseptics and metal salts. The agent is without colour or a different colour from the stain produced on the tooth surface.

Extrinsic tooth discolouration has usually been classified according to its origin, whether metallic or nonmetallic.

- *Non-metallic stains:* The non-metallic extrinsic stains are adsorbed onto tooth surface deposits such as plaque or the acquired pellicle. The possible etiological agents include dietary components, beverages, tobacco...etc.
- *Metallic stains:* Extrinsic staining of teeth may be associated with occupational exposure to metallic salts and with a number of medicines containing metal salts, e.g:
 - The characteristic black staining of teeth in people using iron supplements and iron foundry workers.
 - potassium permanganate producing a violet to black colour when used in mouth rinses;
 - silver nitrate salt used in dentistry causes a grey colour, and
 - stannous fluoride causes a golden brown discolouration.

INTERNALIZED DISCOLORATION

- The stains taken up into the body of enamel or dentine are the same as that causing extrinsic tooth discolouration, including in particular dietary chromogens and the by-products of tobacco smoking. Dental defects permitting the entry of chromogenic material can be classified under the headings of 'developmental and acquired'.
- *1. Developmental defects:* The most important defects are considered under the 'intrinsic tooth discoloration' section. developmental defects may expose dentine either directly or later caused by early loss of enamel as in dentinogenesis imperfecta. Chromogens are then able to enter the dentine directly or facilitated almost certainly by the tubule system.

- *2. Acquired defects:*
- *a) Tooth wear and gingival recession* .Tooth wear is usually considered to be a progressive loss of enamel and dentine due to erosion, abrasion and attrition. As enamel thins the teeth become darker as the colour of dentine becomes more apparent. **Once dentine is exposed the potential of chromogens to enter the body of the tooth is increased.**
- *b) Dental caries:* The various stages of the carious process can be recognized by changes in colour as the disease progresses. For instance, the initial lesion is characterized by an opaque, white spot. The hard, arrested lesion is black having picked up stain from exogenous sources.
- *c) Restorative materials including amalgam:* Some of the materials used in restorative dental treatment may have an effect on the color of teeth. Eugenol and phenolic compounds used during root canal therapy contain pigments which may stain dentine. Some of the poly antibiotic pastes used as root canal medicaments may cause a darkening of the root dentine. Clinicians are familiar with the dark grey to black colour of dentine following the removal of a long-standing amalgam restoration.

HOW CAN WE PREVENT TEETH DISCOLORATION?

- By making a few simple lifestyle changes, you may be able to prevent teeth discoloration. For example, if you are a coffee drinker and/or smoker, consider cutting back or quitting all together. Also, improve your dental hygiene by brushing and flossing regularly and getting your teeth cleaned by a dental hygienist every 6 months.

WHAT TREATMENT OPTIONS ARE AVAILABLE TO WHITEN TEETH?

- Dental treatment of tooth discoloration involves identifying the etiology and implementing therapy.
- A- Diet and habits: Extrinsic staining caused by foods, beverages, or habits (e.g., smoking, chewing tobacco) is treated with a thorough dental prophylaxis and cessation of dietary or other contributory habits to prevent further staining.
- B- Tooth brushing: Effective tooth brushing twice a day with a dentifrice helps to prevent extrinsic staining. Most dentifrices contain an abrasive, a detergent, and an anti tartar agent. In addition, some dentifrices now contain tooth-whitening agents.

- C- Professional tooth cleaning: Some extrinsic stains may be removed with ultrasonic cleaning, rotary polishing with an abrasive prophylactic paste, or air-jet polishing with an abrasive powder. However, these modalities can lead to enamel removal; therefore, their repeated use is undesirable.
- D- Bleaching (tooth whitening): Bleaching includes 2 types of techniques: vital and nonvital.
- **Vital bleaching:** Currently, the bleaching agents most commonly used are carbamide and hydrogen peroxide.
- **In office "power" bleaching** involves the use of a 15-40% **hydrogen peroxide** solution and must be performed by a dental professional because careful isolation of the teeth is required to protect the soft tissues from the caustic effects of the bleaching agent.
- **home bleaching systems** may be used alone or in combination with in-office bleaching. The systems must be used under the careful supervision of dentists or dental hygienists. Patients apply a 10-22% **carbamide peroxide** solution into a custom-made mouth guard. After repeated daily and/or nightly (often while patients sleep) applications for 2-6 weeks, the teeth are gradually bleached.

- Non vital bleaching
- Non vital bleaching is indicated for the treatment of teeth with discoloration secondary to pulpal degeneration. This technique involves placing a mixture of 30% hydrogen peroxide and sodium perborate into the pulp chamber for as long as 1 week.

Thank you

بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ

ETIOLOGY OF PERIODONTAL DISEASES

أعداد:

د. نور صباح أرحيم

Periodontal diseases: ▶

are the most prevalent and multi-factorial diseases that involved hard and soft dental tissues, bacterial colonization, and immune responses of the host. periodontal diseases affecting a large population worldwide. plaque , being the primary etiological agent, a genetic origin may be play a part. there are two types of periodontal diseases, gingivitis and periodontitis.

GINGIVITIS:

It is inflammation of the gingiva in which the junctional epithelium remain attached to the tooth at its original level. It is characterized by areas of redness and swelling, and there is a tendency for the gingiva to bleed easily. Gingivitis is limited to the epithelium and gingival connective tissues. It is important to note that there is no tissue recession or loss of connective tissue or bone. The occurrence of gingivitis is wide spread in the population. It is reversible condition .



Periodontitis: ▶

it is a common , chronic inflammatory disease (inflammation of the supporting tissues of the teeth leading to permanent destruction of these tissues) caused by the accumulation of bacterial matrix at the gingival line. it is characterized by clinical attachment loss, periodontal pocketing and alveolar bone loss. it is irreversible disease can cause a breakdown of the periodontium result in the loss of the tissue attachment and destruction of the alveolar bone.



Contained gingivitis:

in some individuals gingivitis may not progress to periodontitis for a long period of time even if is not treated and this condition depend on host response and the pathogenicity of bacteria.

Risk factors:

Can be defined as characteristics or factors that when present increases the risk that an individuals will get the diseases. It is important to make the distinction that risk factors are associated with a disease but do not necessarily cause the disease, can be classified into local and systemic risk factors.

LOCAL RISK FACTORS:

1-anatomical risk factor

- enamel pearls.
- root groove (palatogingival groove).
- furcation.
- gingival recession.

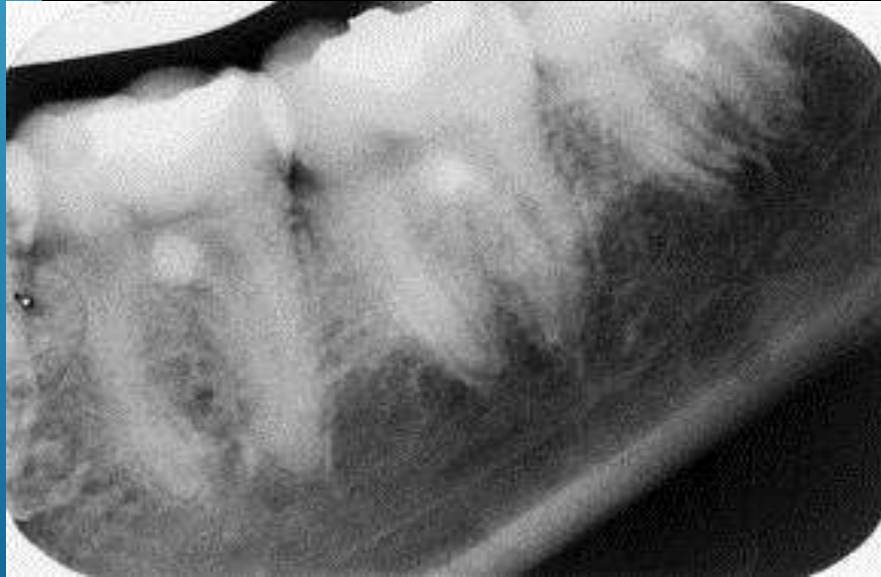
2-tooth position

- misalignment.
- crowding.
- tipping.
- migration.
- occlusal forces.

3-Iatrogenic risk factor

- partial dentures
- overhang restoration.
- orthodontic appliance.

ENAMEL FLAKES



PALATOGINGIVAL GROOVE

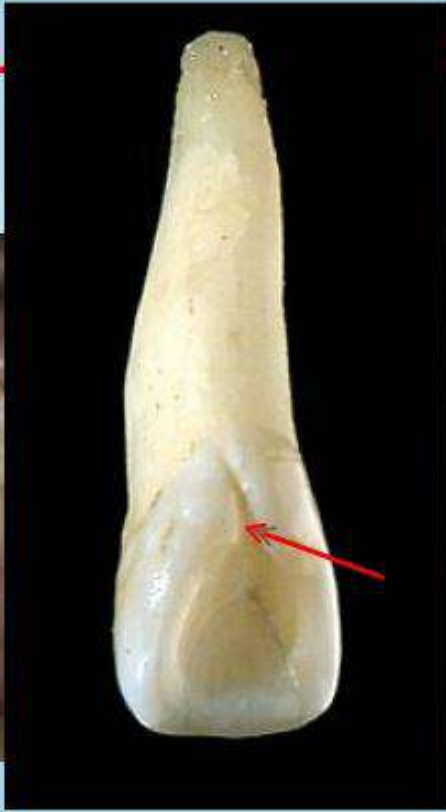
MAXILLAR INCISORS

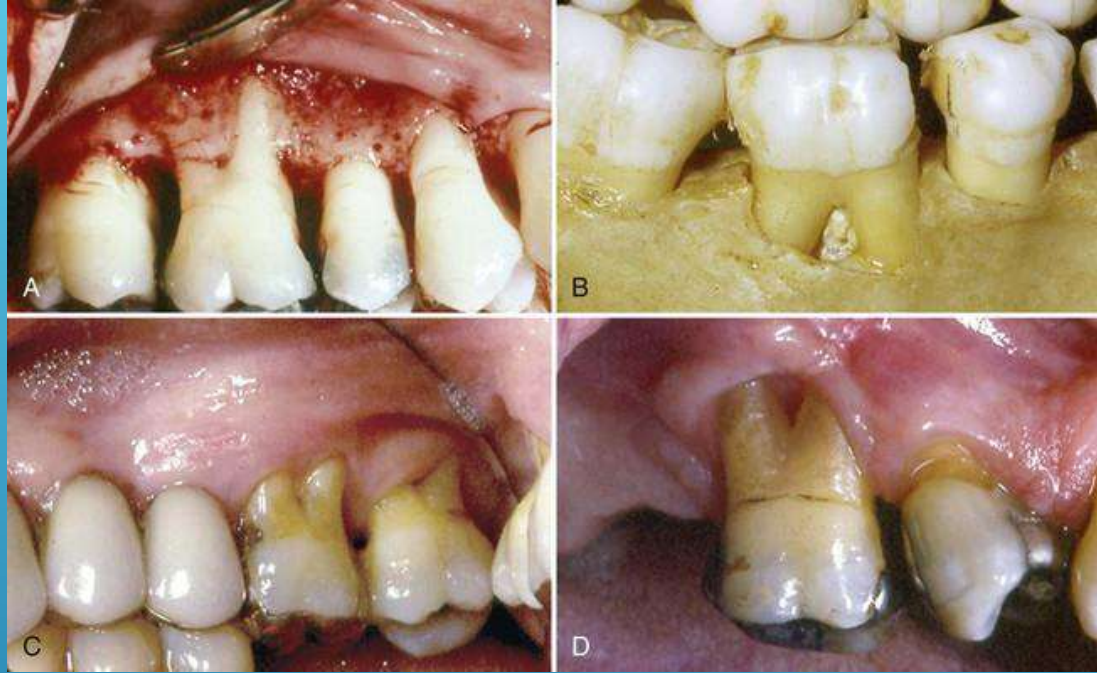
Variations
Peg shaped lateral
and small

The missing

Large
cingulum

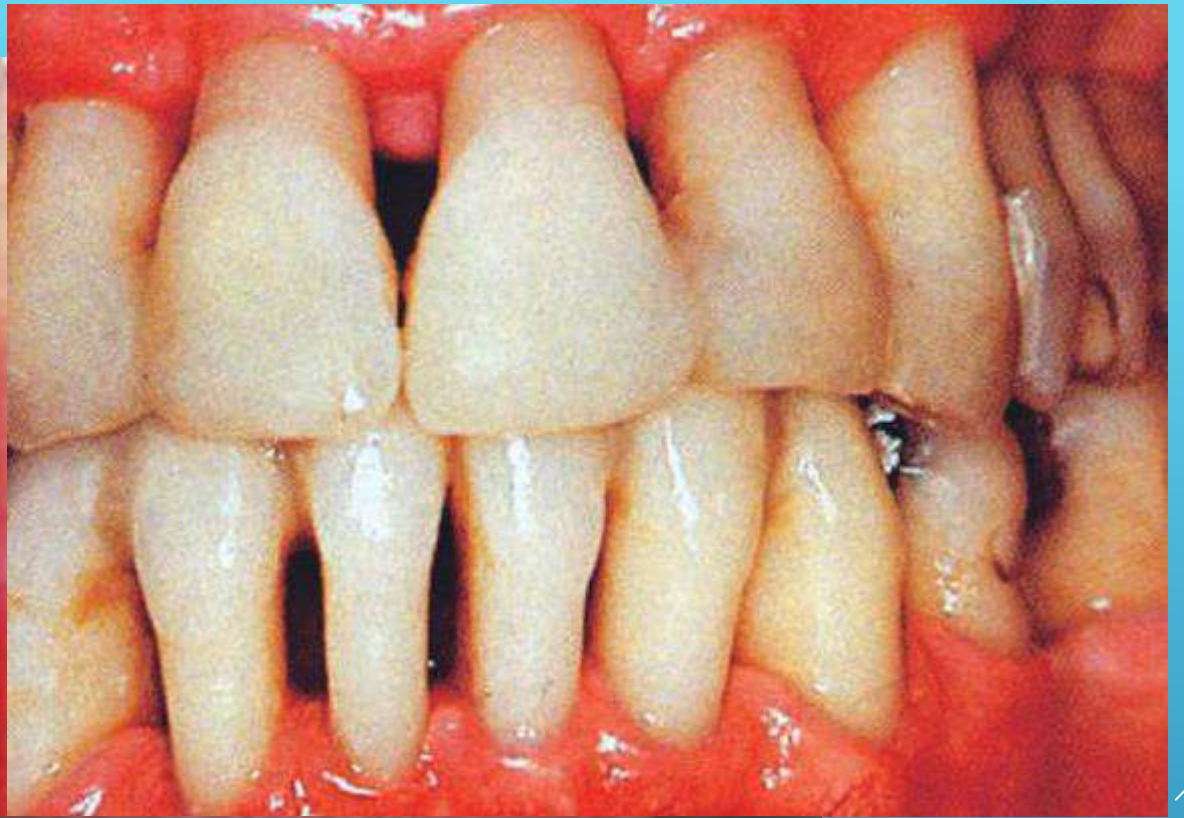
Deep
distal

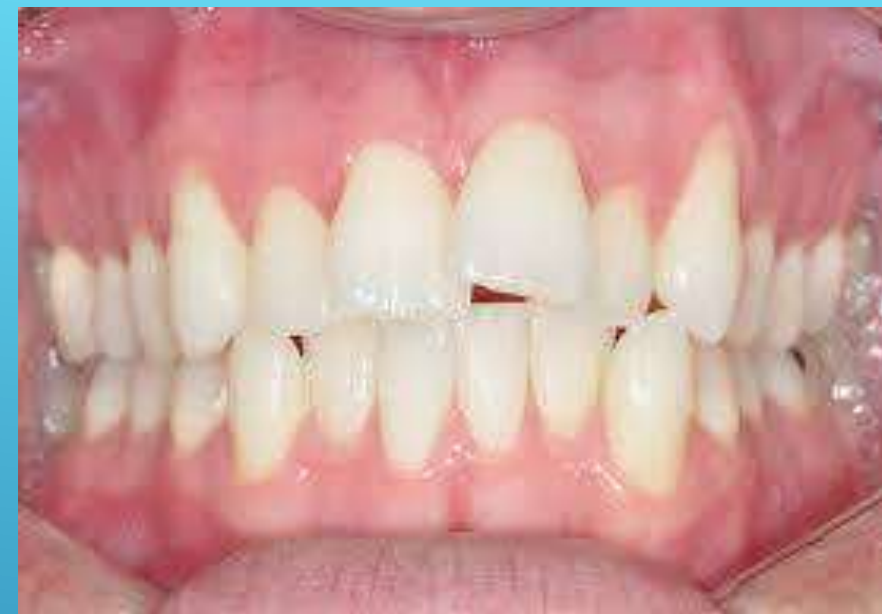


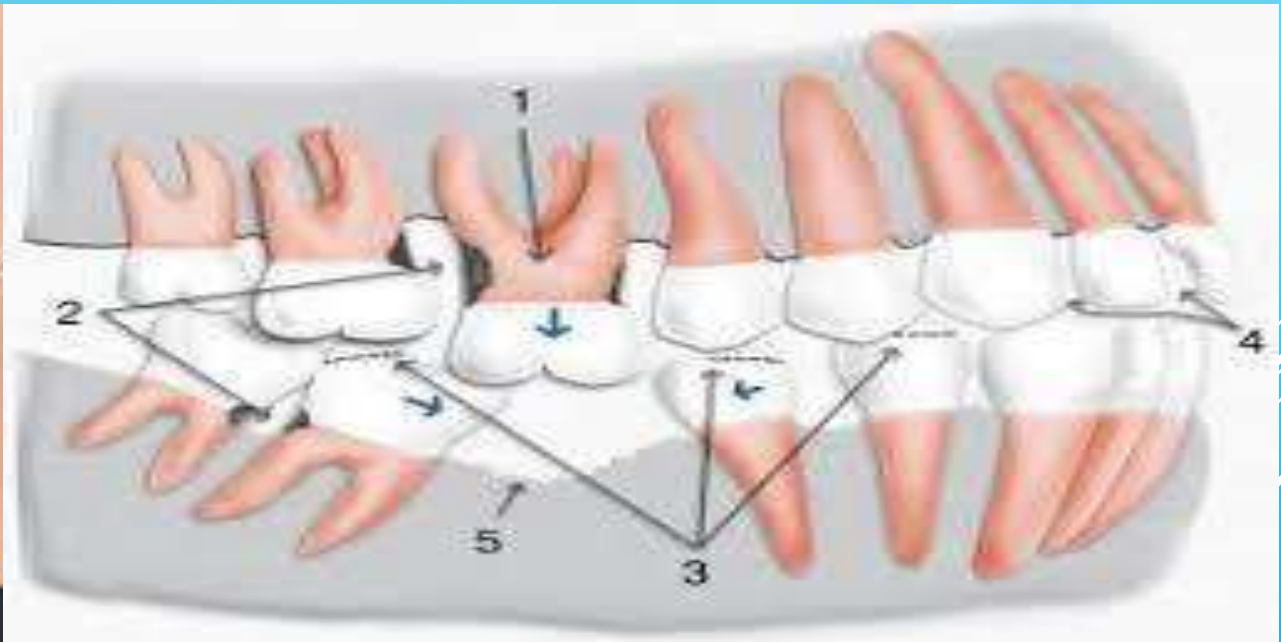


FURCATION AREA

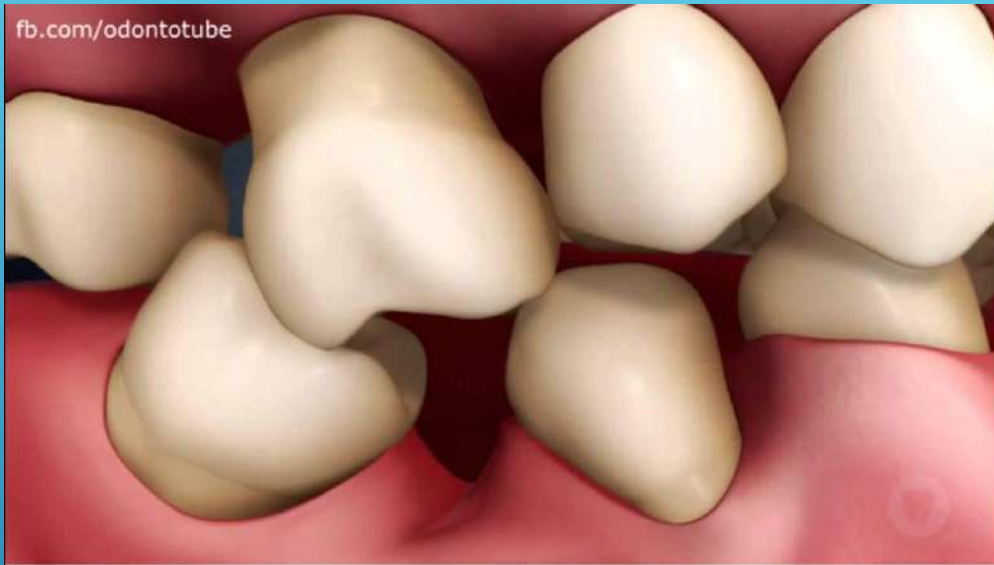


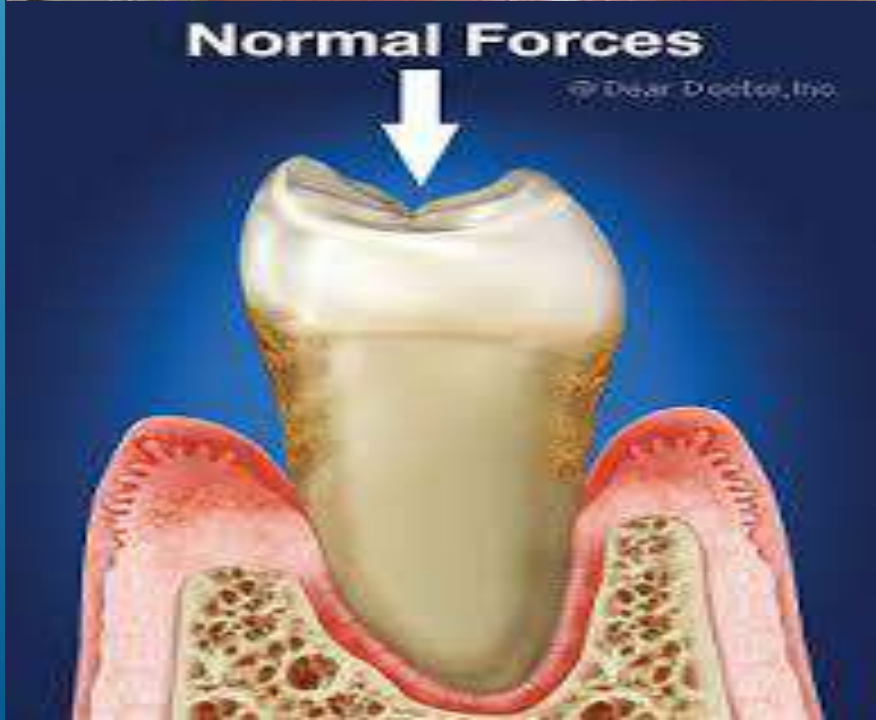




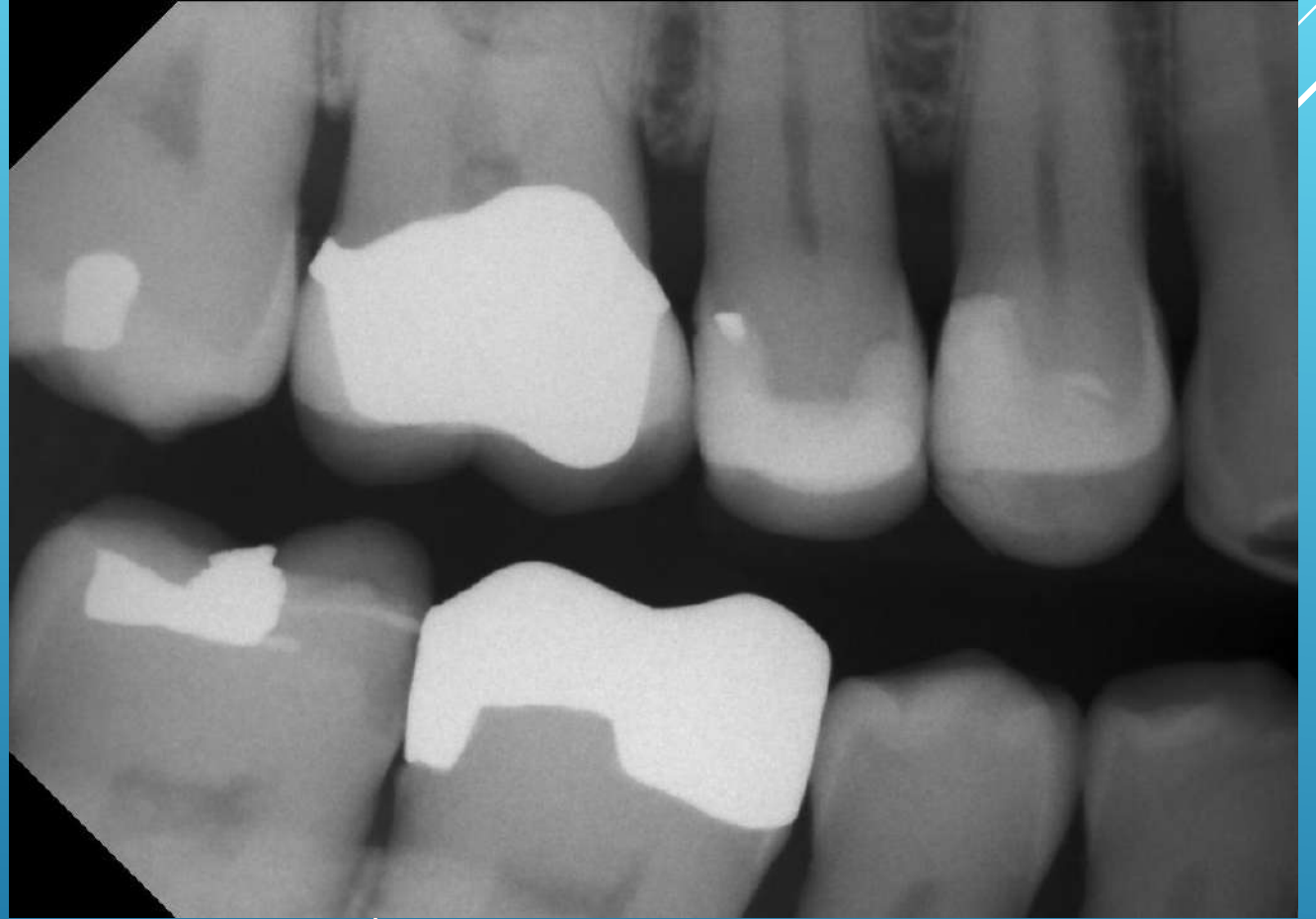


fb.com/odontotube





overhang restoration







partial denture



Orthodontic appliance

THANK YOU



GINGIVA 2

PRESENTED BY :

DR. NOOR SABAH IRHAYYIM

Normal microscopic feature of gingiva

- The gingiva consists of fibrous connective tissue known as lamina propria covered by stratified squamous epithelium. ➤
- gingival epithelium may be differentiated as follows: ➤
 - 1-oral epithelium:** which faces the oral cavity ➤
 - 2-Sulcular epithelium:** which faces the tooth in the gingival sulcus without being in contact with the tooth surface. ➤
 - 3-Junctional epithelium:** which provides the contact between the gingiva and the tooth. ➤

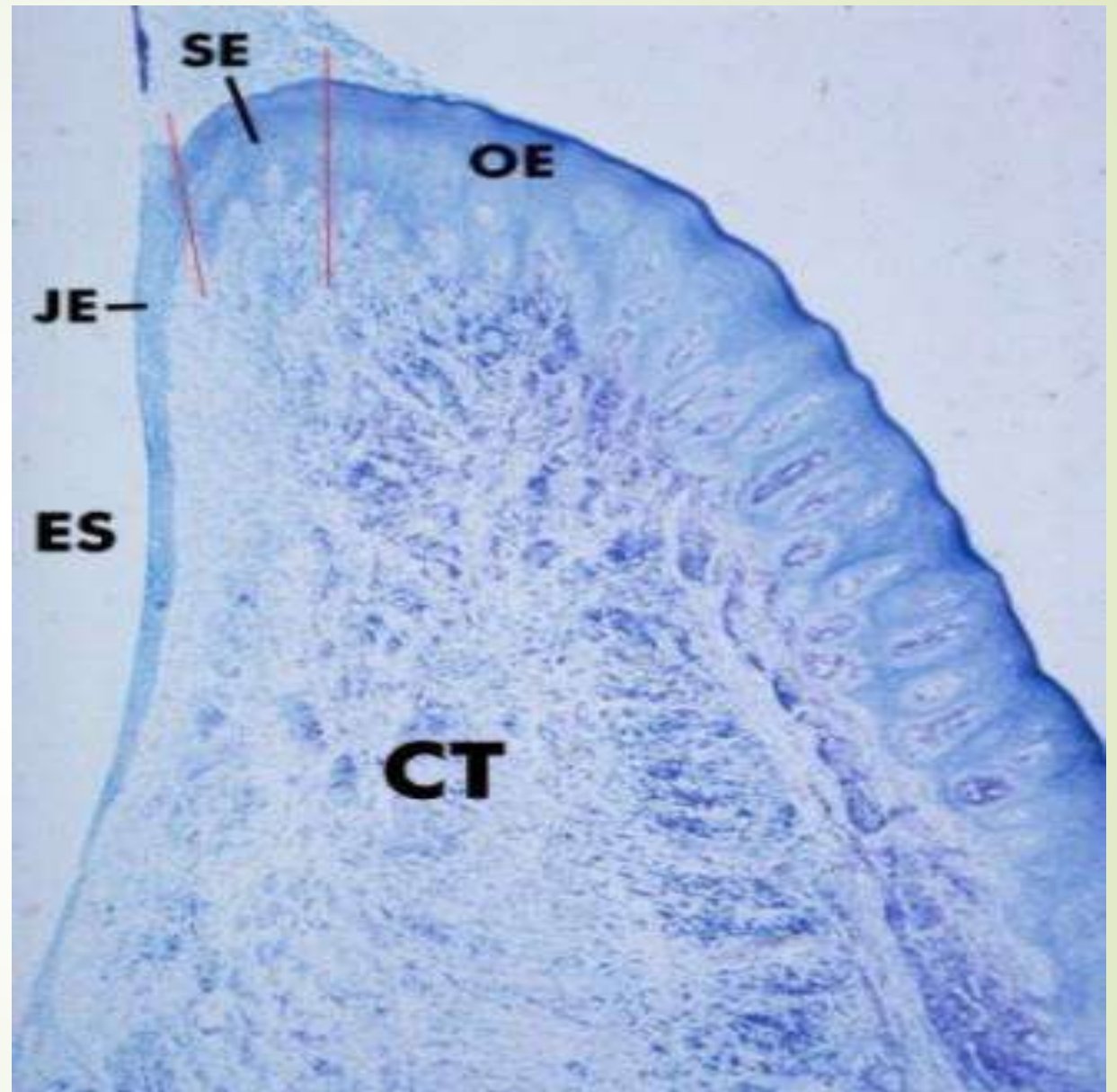
CT, gingival
connective
tissue

ES, ⑥ ⌚
enamel space

JE, ⑥ ⌚
junctional
epithelium

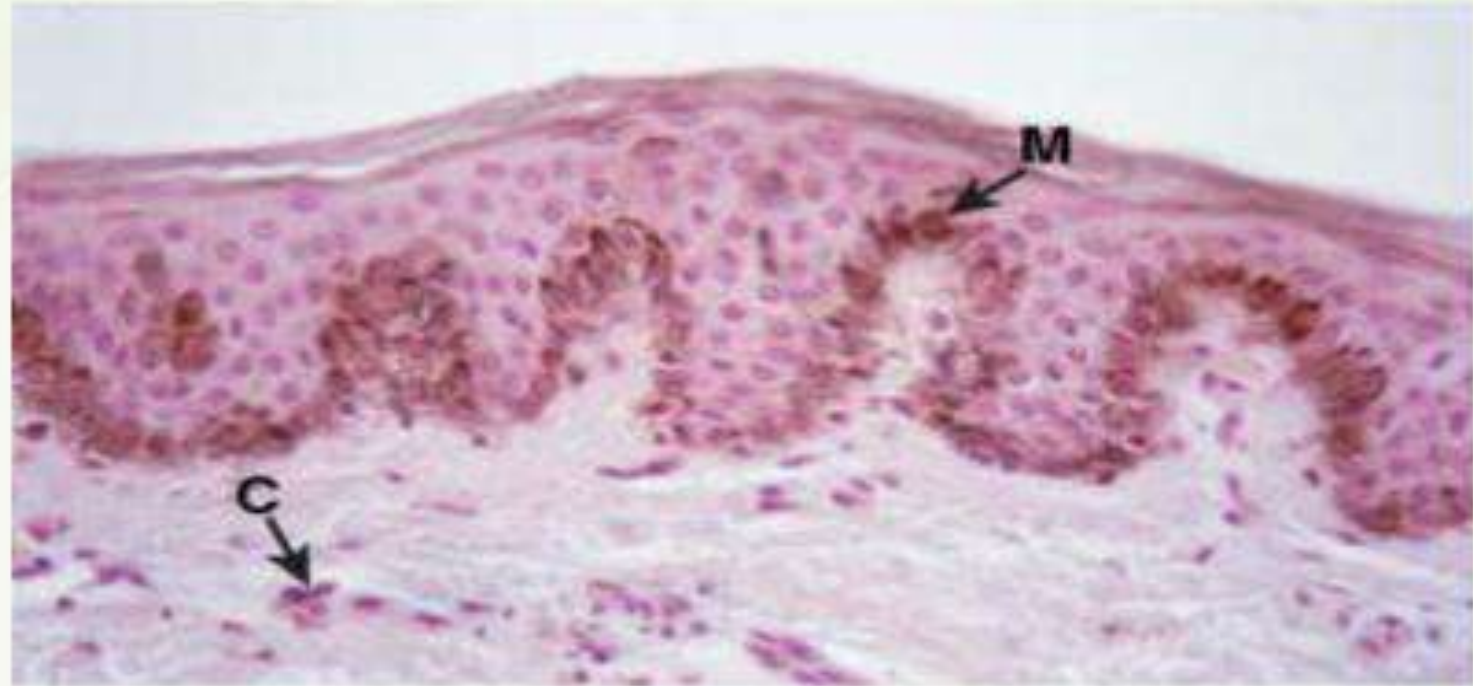
OE, oral ⑥ ⌚
epithelium

SE, ⑥ ⌚
sulcular
epithelium



Oral epithelium:

- It covers the crest and the outer surface of the marginal and attached gingiva. It is either keratinized (no nuclei) or parakeratinized (retained nuclei). The boundary between the oral epithelium and the underlying connective tissues has a wavy course. ➡
- The projection of the epithelium cells into the connective tissues are known as (**Rete Pegs**). The intervening connective tissue portion which project into the epithelium are called connective tissue papillae. ➡
- This alternating pattern of depression and protuberances of the connective tissue papillae and epithelium rete pegs thought to give the attached gingiva (the stippling appearances) ➡



Copyright © 2006 by Saunders, an imprint of Elsevier Inc.

Pigmented gingiva of dog showing melanocytes (M) in the basalepithelial layer and melanophores (C) in the connective tissue (Glucksman technique).

The oral epithelium consists of four layers of cells

1-Stratum basale: Basal layer of cuboidal cells along the basement membrane . This is where epithelial cell replication and cell differentiation begins.(melanocytes are found in this layer) ➤

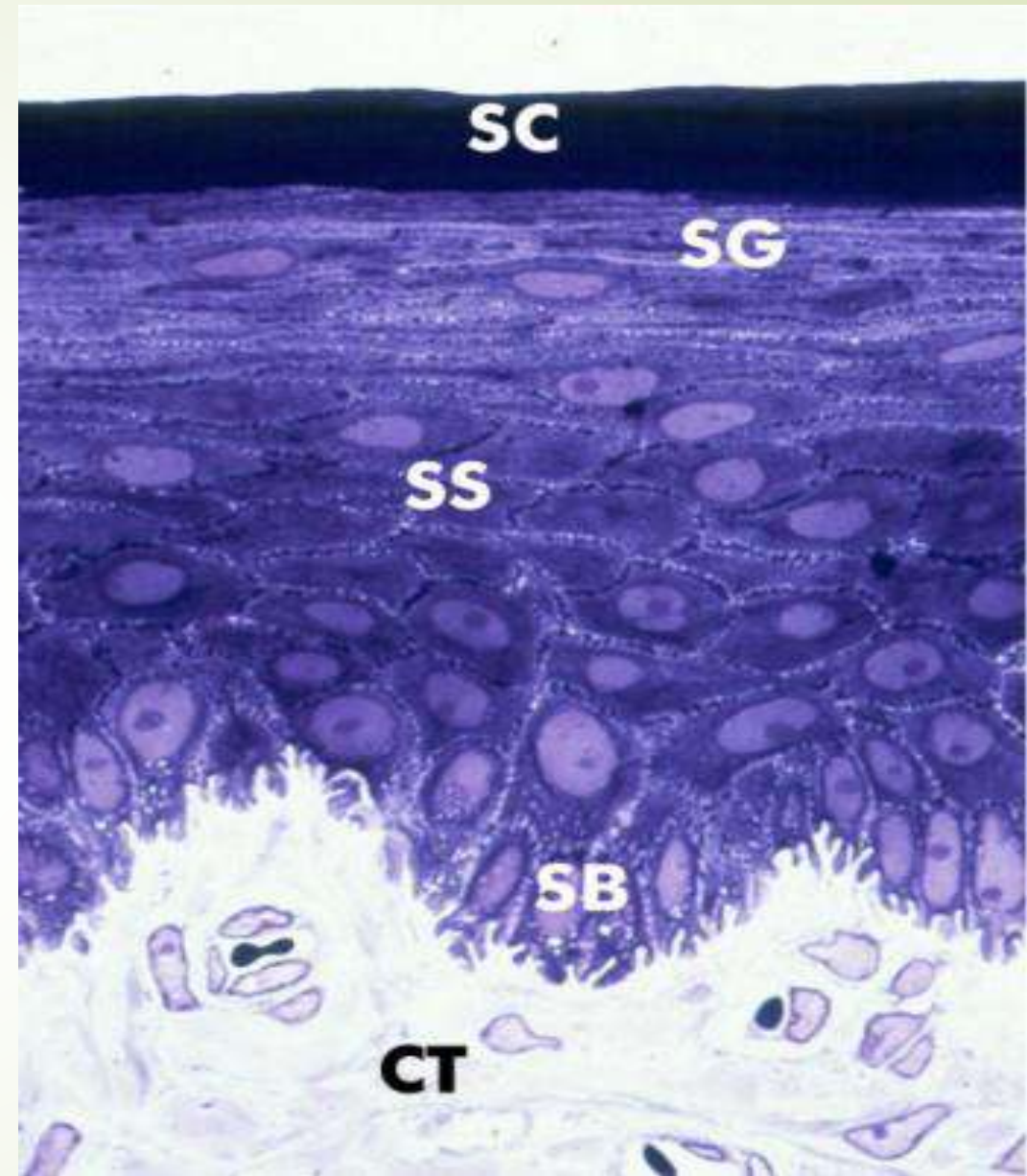
2-Stratum spinosum: the cells appear to have cytoplasmic spines. This is the thickest cell layer and Langerhans cells are found in this layer. ➤

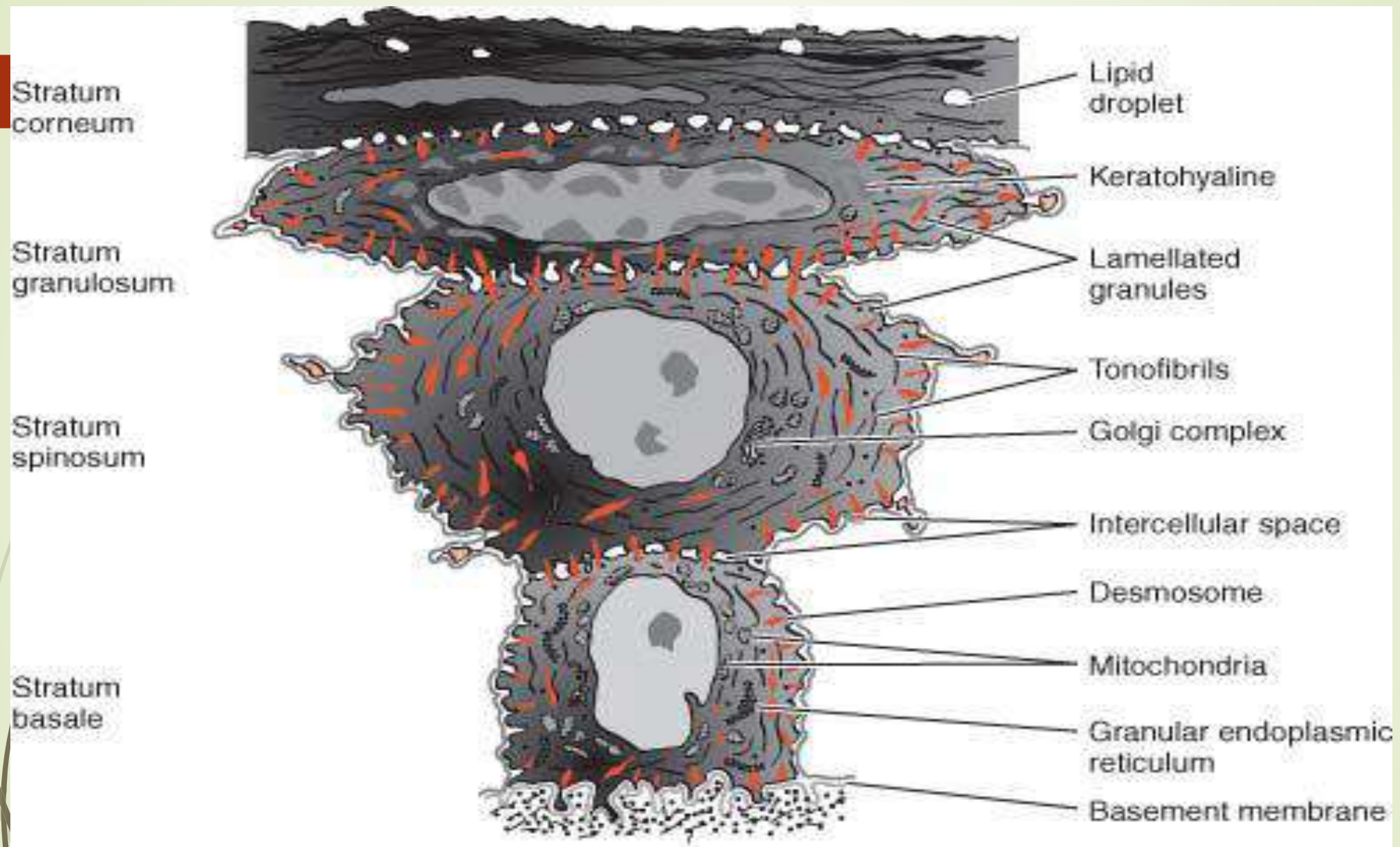
3-Stratum granulosum: Keratohyaline granules may be seen in this layer. Cells appear to be flattened. ➤

4-Stratum corneum: This is the layer where both para or the orthokeratinized occur. It is the most superficial layer. ➤

-The epithelium cells are formed as basal cells and gradually they undergo the process of keratinization, this is achieved by proliferation and differentiation of these cells (change to the characteristic of each of the cell layer) as they migrate towards the surface layer. ➤

SC, stratum corneum
(cornified layer)
SG, stratum
granulosum (granular
layer)
⌚Ⓜ **SS**, stratum
spinosum (spinous
layer)
⌚Ⓜ **SB**, stratum basale
(basal layer)
CT, connective Ⓜ⌚
tissue

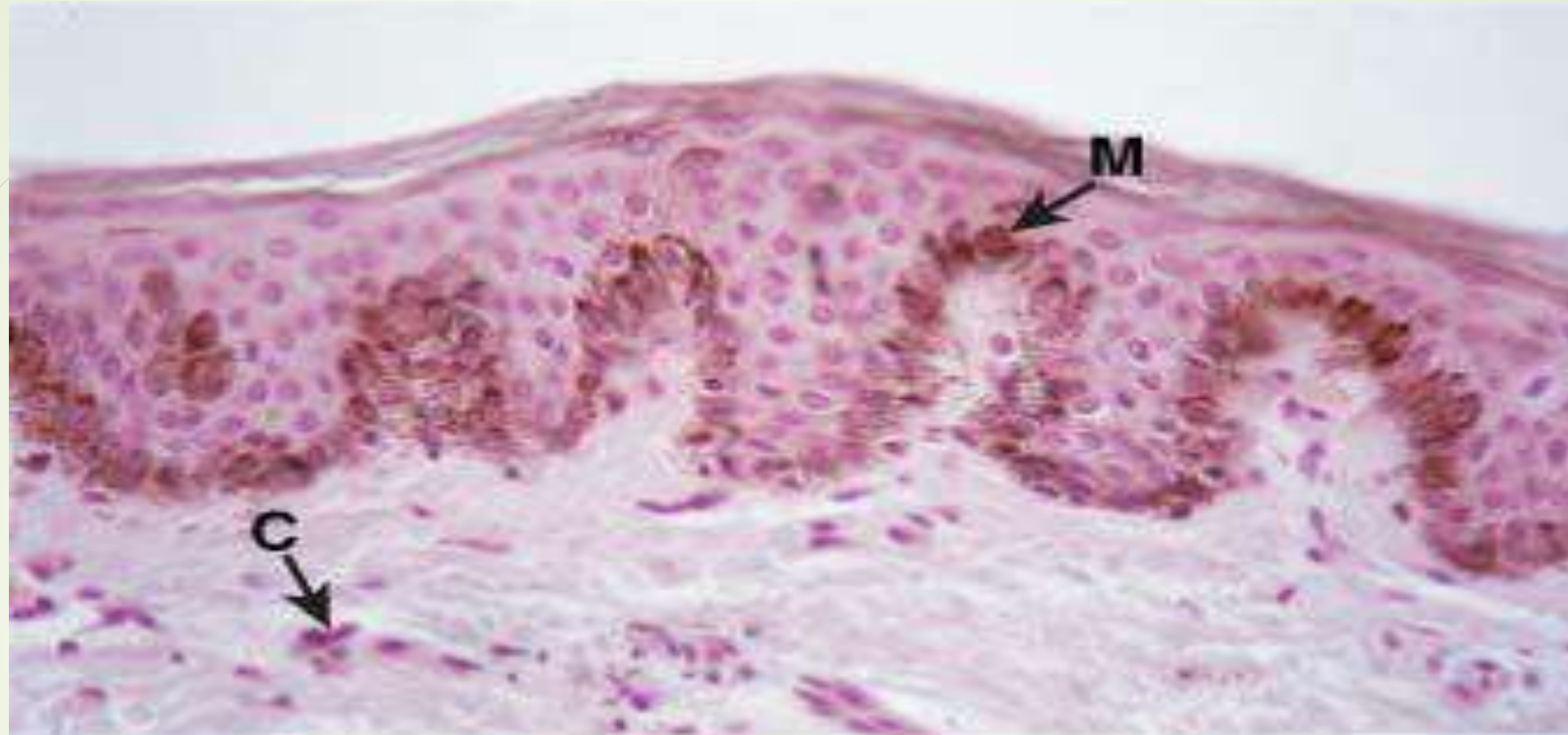




Copyright © 2006 by Saunders, an imprint of Elsevier Inc.

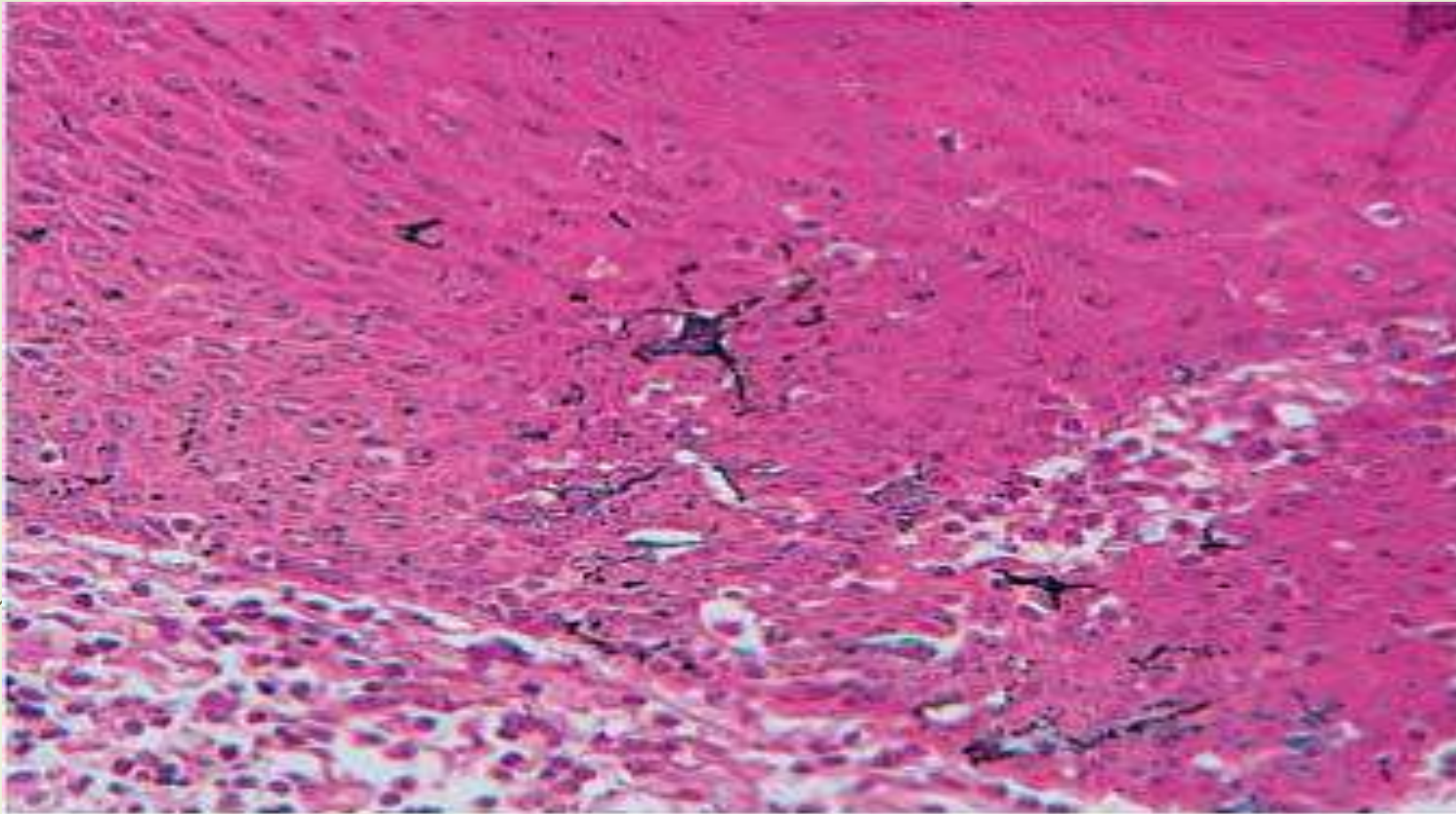
The oral epithelium contains the following types of cells:

- 1- Keratinocytes cells:** These are Keratin producing cells which comprise about **90%** of the total cell population. These cells undergo continuous proliferation and differentiation from basal layer to the surface of epithelium. Keratin may be found in the stratum corneum and contribute to the protective function of epithelium.
- 2-Melanocytes:** cells of basal layer that produce melanin pigment granules.
- 3-Langerhans cells:** these cells play a role in the defense mechanisms of the oral epithelium. They have an immunological function by recognizing and processing antigens.
- 4-Merkel cells:** these are located in the deeper layers of the epithelium, harbor nerve ending. They have been identified as tactile receptors.



Copyright © 2006 by Saunders, an imprint of Elsevier Inc.

Pigmented gingiva of dog showing melanocytes (M) in the basalepithelial layer and melanophores (C) in the connective tissue (Glucksman technique).



Copyright © 2006 by Saunders, an imprint of Elsevier Inc.

Human gingival epithelium, oral aspect. Immunoperoxidase technique showing **Langerhans cells**. Function: On infection of an area of skin, the local Langerhans' cells will take up and process microbial antigen to become fully-functional antigen-presenting cells

Under a normal condition, there is complete equilibrium between cell renewal and desquamated (cell turn over). It takes approximately 3-4 weeks for keratinocytes to migrate from basal layer until reach the outer epithelial surface, where it become desquamated from stratum corneum. ➤

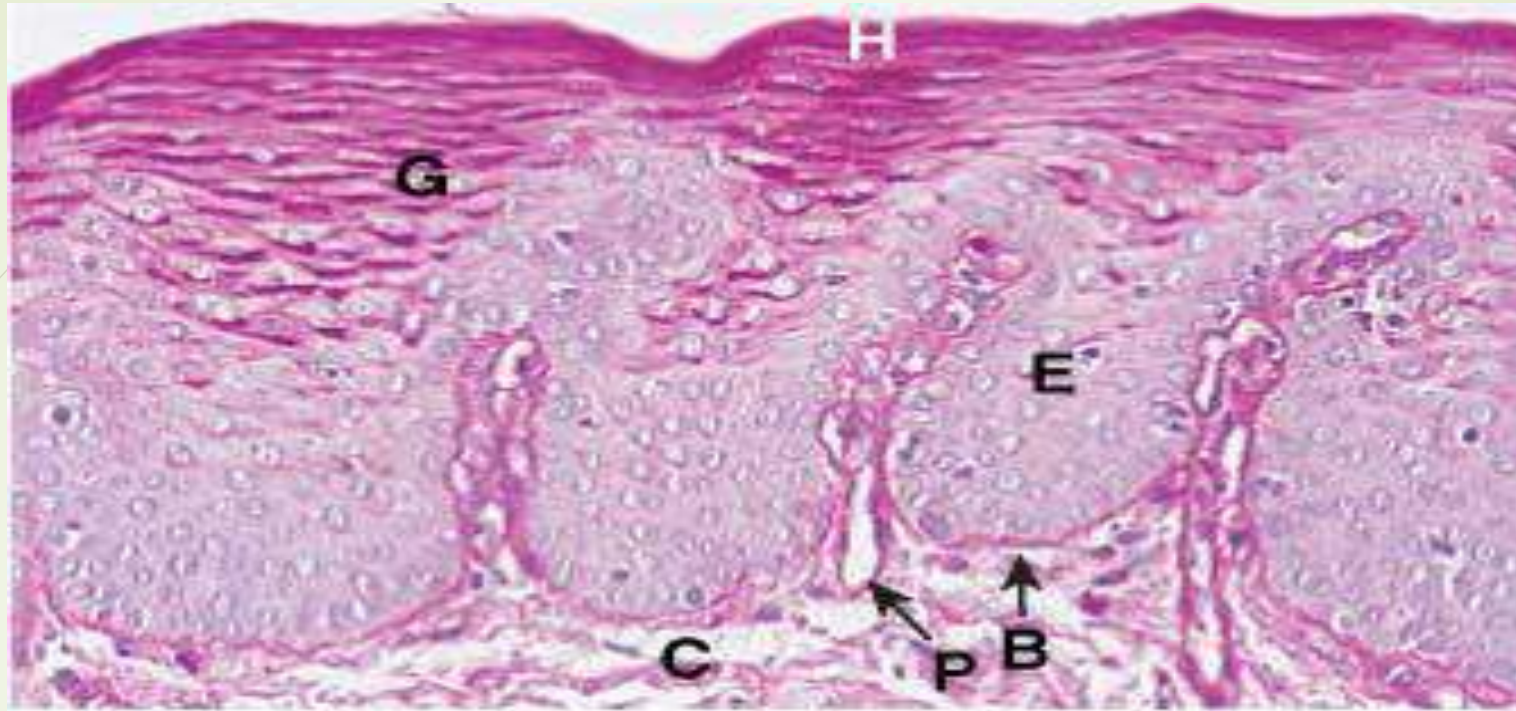
Basal cells are found immediately adjacent to the connective tissue and are separated from this tissue by a basement membrane (basal lamina). ➤

The basement membrane consists of : ➤

1-Lamina lucida: which is located immediately beneath the basal cell layer. ➤

2- lamina densa: located beneath the lamina lucida, from this structure the anchoring fibers project into the connective tissue. ➤

-The epithelial cell are joined together by structure known as desmosome, which is composed of two hemi desmosome separated from each other by granulated material. ➤



Copyright © 2006 by Saunders, an imprint of Elsevier Inc.

Normal human gingiva stained with the periodic acid-Schiff (PAS) histochemical method. The basement membrane (B) is seen between the epithelium (E) and the underlying connective tissue (C). In the epithelium, glycoprotein material occurs in cells and cell membranes of the superficial hornified (H) and underlying granular layers (G). The connective tissue presents a diffuse, amorphous ground substance and collagen fibers. The blood vessel walls stand out clearly in the papillary projections of the connective tissue (P).



A hemi desmosome composed from the following structures:

- 1- the outer leaflets (OL) ➡**
- 2-the inner leaflet (IL) ➡**
- 3-the attachment plaque (AP) ➡**

Sulcular Epithelium

- 1-It lines the gingival sulcus . ➡
- 2- It is a thin, non keratinized stratified squamous epithelium without rete ➡ pegs and extend from the coronal limit of the junctional epithelium to the crest of the gingival margin.
- 3-The sulcular epithelium important because it is thin and may act as a ➡ semipermeable membrane through which tissue fluid from the gingiva seep into the sulcus and makes easier for bacterial products of dental plaque to penetrate into the connective tissue of the gingiva and stimulate the inflammation and tissue destruction , **That why the sulcular epithelium id considered as a poor barrier against bacterial infection.**

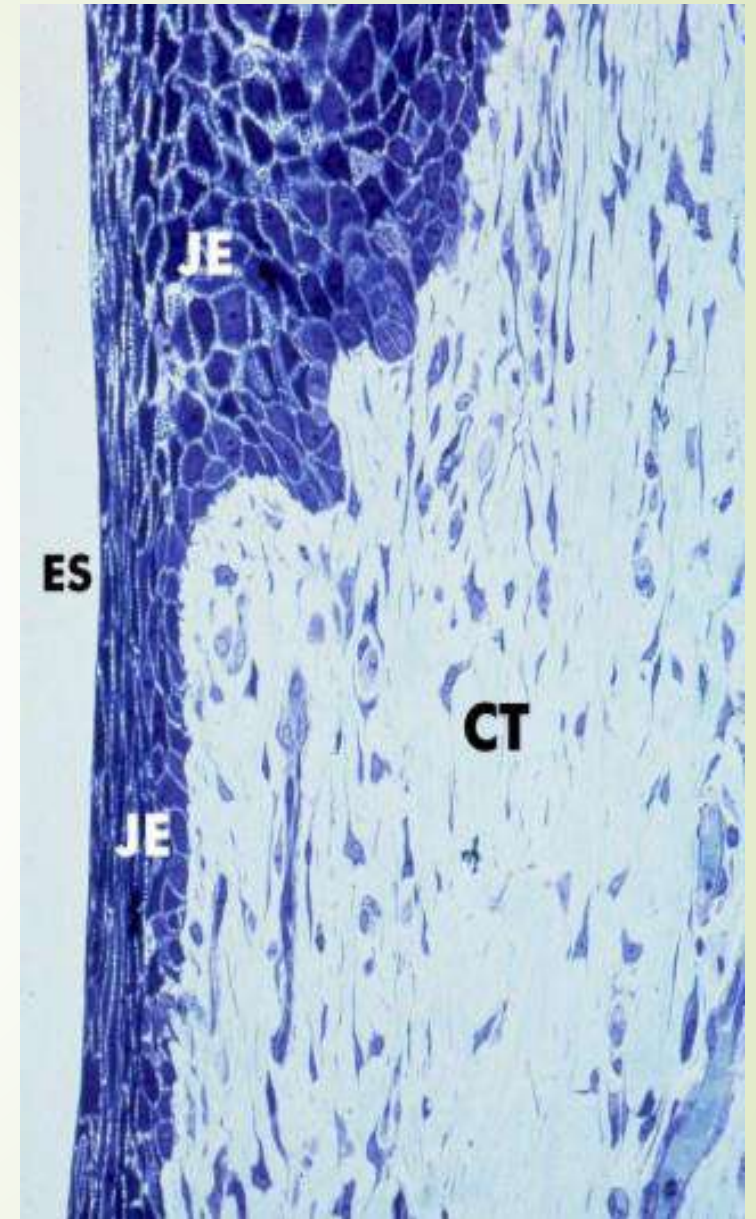
Junctional epithelium (JE)

- the epithelium that attaches the gingiva to the surface of the tooth ▶
- it consists of stratified squamous non-keratinized epithelium ▶
- it is 3-4 layers thick in early life but the numbers of increases with age to 10-20 ▶
- it is thicker in coronal portion but become thinner towards cemento-enamel junction only a few cell layers ▶
- the junctional epithelium cells can be grouped in two layers: the basal and suprabasal layer. ▶

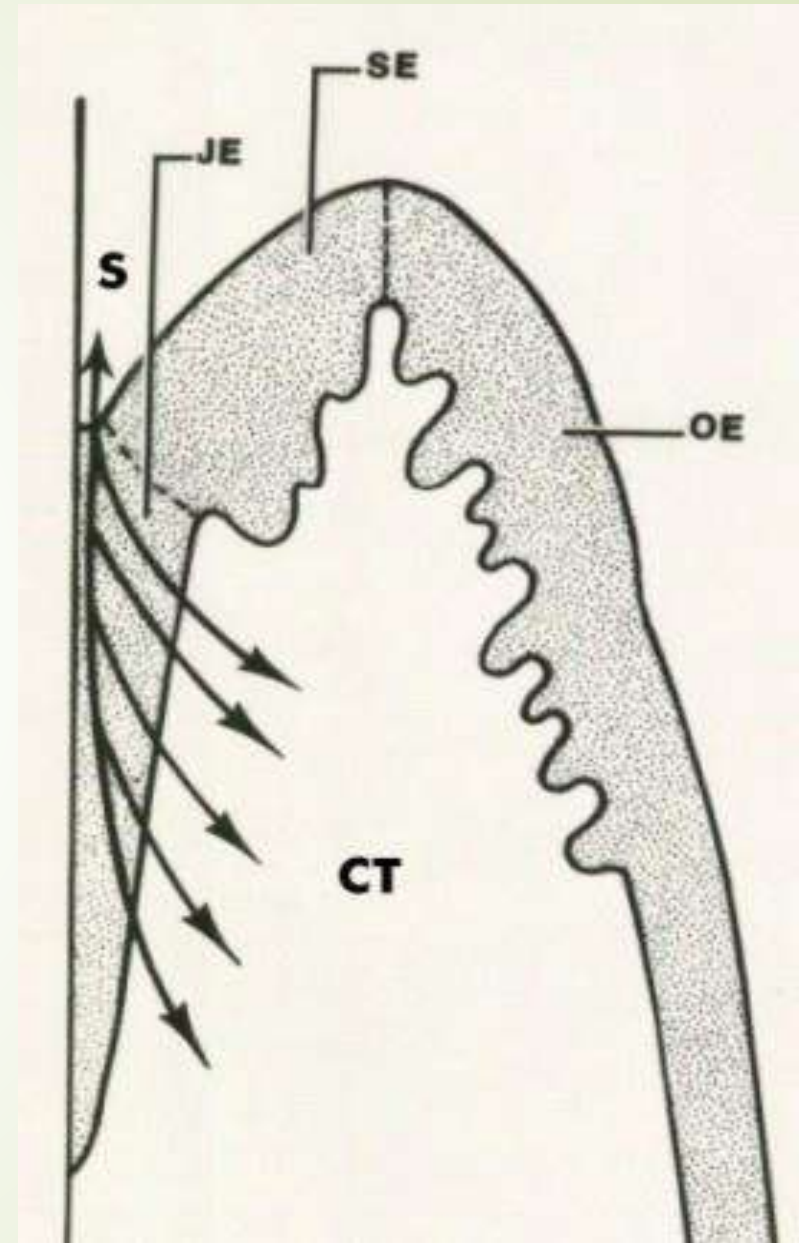
The epithelium is continuously renewed through cell division in the basal layer and the cell migrate coronally to the base of the gingival sulcus from they are shed (cell turn over) ▶

- the JE assume a key role in maintenance of periodontal health, it creates the firm epithelium attachment that connect the soft tissue to the tooth surface, it is quite permeable and thus serve as a pathway for diffusion of products plaque bacteria to connective tissue, there is also diffusion in the opposite direction moving towards the sulcus of host defense substance, this help to maintain the immune response. ▶

CT, connective tissue
ES, enamel space
JE, junctional epithelium



CT, connective
tissue
JE, junctional
epithelium
OE, oral
epithelium
S, gingival sulcus
SE, sulcular
epithelium



There are distinct differences between the JE, sulcular and the oral epithelium :

- 1-The size of the cells in the JE is relatively larger than the oral epithelium ➤
- 2-the intercellular space in the JE is wider than in the oral epithelium. The intercellular space of the JE is preferred route for tissue fluid and inflammatory cells to migrate from the CT to gingival sulcus. ➤
- 3-The number of desmosome (intercellular junction) is fewer in the junctional epithelium than in the oral epithelium , this may explain the JE susceptibility to tear during probing and it is greater permeability to migrate cells and fluid. ➤
- 4-The sulcular and junctional epithelium are not as thick as the oral epithelium, because they are not keratinized and in health have no retepegs. ➤
- 5-JE turnover rate is very high (4-6) days compared to oral epithelium that has longest turn over rate (6-12 days) or (up to 40 days). ➤
- 6-JE forms the attachment of the gingiva to the tooth surface while oral and sulcular epithelium have no attachment to the tooth surface. ➤

Gingival connective tissue:

The connective tissue of the gingiva known as the lamina propria and consists of two layers:

1-The papillary layer: it consists of papillary projections between the epithelial rete pegs.

2-The reticular layer: it is contiguous with the periosteum of the alveolar bone

-The major components of the CT are:

1-collagen fibers 60%

2-cells 5%

3-ground substances, blood vessels, nerve, and lymphatics(35%).



Cells of gingival CT:

1-Fibroblasts ➡

2-Mast cells ➡

3-macropgage ➡

4-inflammatory cells ➡



Ground substances of gingival CT

- 1-the matrix fill the space between fibers, cells and have high content of water, electrolytes, nutrition, metabolites, glycoprotein and proteoglycan.
- 2-it is produced by fibroblast and some from the mast cells and other from the blood.
- 3-it is a medium for the cells of CT to embedded and for maintained the normal function of the CT

Blood vessels of gingiva

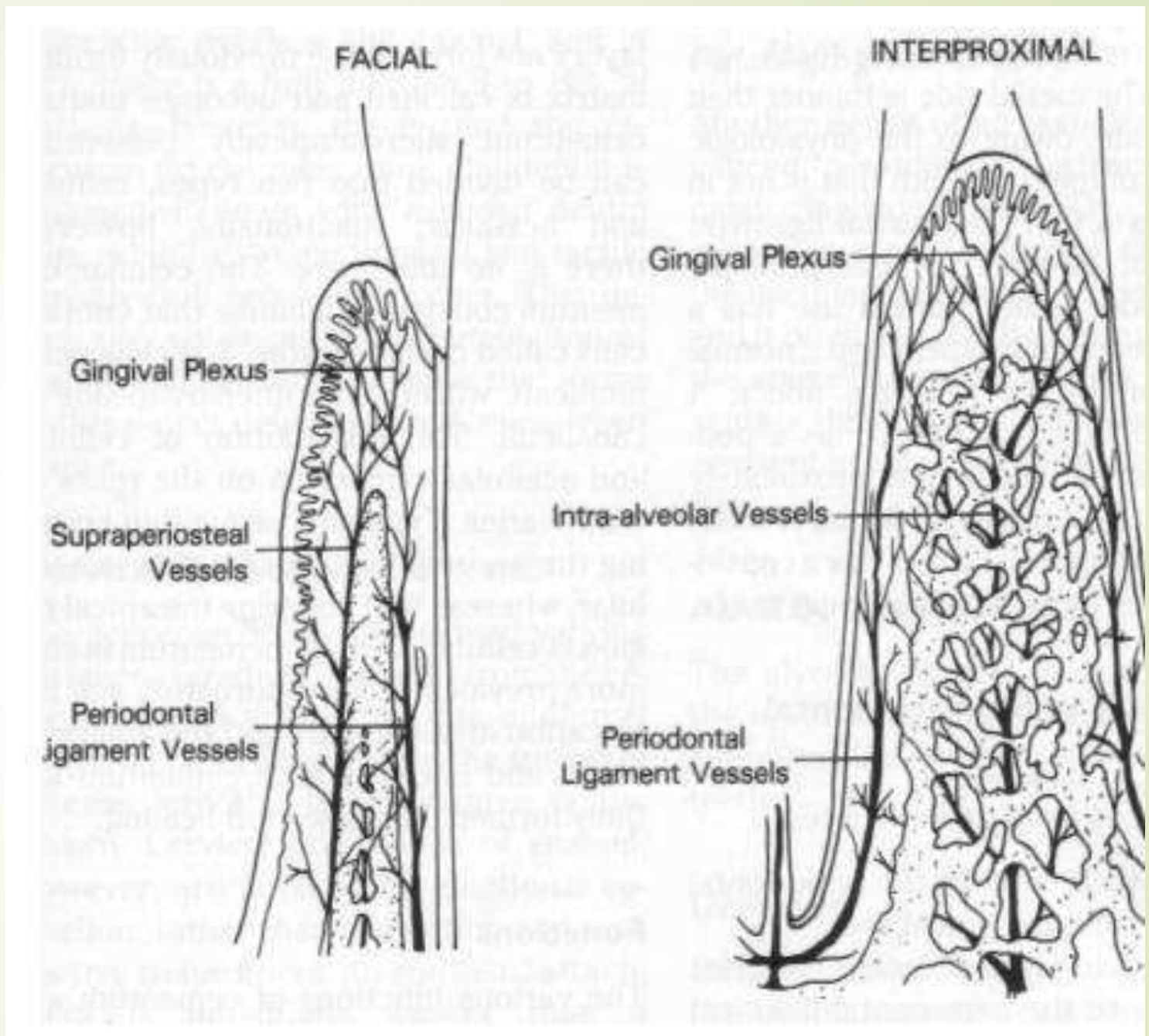
- gingival tissue has rich vascular supply which arise from the terminal branches of internal maxillary artery.

- consist of

1-supra periosteal vessels'

2-vessels from the PDL and bone








Innervation of the gingiva

Nerve supply to gingiva derived from the terminal branches of the maxillary and mandibular branches of the trigeminal nerve. ➤





THANK



YOU

Pathogenesis of periodontal disease inflammatory responses in the periodontium

م. سها أسود دهش العزاوي

B.D.S, MSc. Periodontology

Periodontal disease is initiated and sustained by factors (substances) produced by the subgingival microbiota (the biofilm). Some of these substances can **directly** injure host cells and tissues. Other microbial constituents may **activate inflammatory** or cellular and humoral **immune systems** that cause damage to the periodontal tissues. It is the latter pathway which accounts for most injury to the periodontal tissues.

Microbial virulence factors

- *Lipopolysaccharide*
- *Bacterial enzymes*
- *Microbial invasion*
- *Fimbriae*
- *Bacterial DNA*

Host-Derived Inflammatory Mediators

- *Cytokines*
- *Prostaglandins*
- *Matrix metalloproteinases*

Lipopolysaccharide

- large molecules composed of a lipid component (lipid A) and a polysaccharide component. They are found in the outer membrane of gram-negative bacteria (LPS is frequently referred to as endotoxin). **They elicit strong immune responses in animals.**
- TLR-4 recognizes LPS from gram-negative bacteria.
- LPS triggers a series of intracellular events, the net results of which are the increased production of inflammatory mediators (most notably cytokines) and the differentiation of immune cells (e.g., dendritic cells) for the development of effective immune responses against the pathogens. **LPS is of key importance for initiating and sustaining inflammatory responses in the gingival and periodontal tissues.**
- Porphyromonas gingivalis has an atypical form of LPS that is recognized by both TLR-2 and TLR-4.

Microbial invasion

The invasion process is considered a common strategy shared between various microbial pathogens and is thought to have evolved in order to access nutrients and shelter from host defences. *P. gingivalis* has the ability to bind and subsequently invade a range of eukaryotic cells other than oral epithelial cells, including fibroblasts, endothelial cells, and multiple cell lines, although it is not clear whether the mechanisms involved are the same for all cell types. The establishment of microbial species within an intracellular environment represents a challenging issue for extracellular and intracellular host defence mechanisms. The intracellular environment is not exposed to normal host immune defense mechanisms or many therapeutic agents, which represents a feature of a number of virulent organisms. It has been shown that *Aggregatebacter actinomycetemcomitans* can penetrate human epithelial cells by switching its morphological phenotype during the invasion process.

Enzymes and Noxious Products

Microorganisms produce a variety of soluble **enzymes** that may digest extracellular host proteins and other molecules and thereby produce nutrients for bacterial growth. In addition to enzymes, bacteria also release numerous, **harmful metabolic waste products**, such as ammonia, indole, hydrogen sulfide, and butyric acid. Amongst the enzymes released by bacteria in the biofilm, proteases (proteinases) are capable of digesting collagen, elastin, fibronectin, fibrin, and various other components of the intercellular matrix of both epithelial and connective tissues

Proteinases (proteases)

Periodontal disease results in tissue degradation, and thus **proteases, derived both from the host and from bacteria, are central to the disease processes.** Proteinases (collagenase, elastase-like and trypsin-like, as well as serine and cysteine proteinases) cleave proteins by hydrolyzing peptide bonds and may be classified into two major classes, endopeptidases and exopeptidases, depending on the location of activity of the enzyme on its substrate.

Proteinase inhibitors

Release of proteinases in the gingiva and the crevicular area promotes inflammatory reactions and contributes to connective tissue damage via several pathways. In contrast, proteinase inhibitors would dampen the inflammatory process. Among such inhibitors **alpha-2 macroglobulin (A2-M)** and **alpha1 antitrypsin (A1-AT)** must be recognized. In fact, **gingival collagenase inhibition by A2-M** has been demonstrated to occur in gingival tissues and **polymorphonuclear leukocyte (PMN) collagenase is inhibited by A1-AT**. Many host and microbial enzymes are likely to be present in the crevice at any one time. Realizing the potentially destructive features of such enzymes, consideration should be given to the source of these enzymes, their relative proportions and the inhibitory mechanisms available within the crevice. **The main enzyme activity is host derived and specific and non-specific inhibitors are plentiful within the crevice and thus enzyme activity will be localized and shortlived**

Matrix metalloproteinases (MMP)

Matrix metalloproteinases (MMPs) hydrolyses components of the extracellular matrix. These proteinases play a central role in many biological processes, such as embryogenesis, normal tissue remodeling, wound healing, and angiogenesis, and in diseases such as atheroma, arthritis, cancer, and tissue ulceration. Currently **28 MMP genes have been identified** in humans. The periodontium is structurally comprised of fibrous elements, including collagen, elastin, and glycoproteins (laminin, fibronectin, proteoglycans), minerals, lipids, water, and tissue-bound growth factors. In addition there are variety of extracellular matrix components, including tropocollagen, proteoglycans, and other proteins (elastin, osteocalcin, osteopontin, bone sialoprotein, osteonectin, and tenascin). All of these matrix components are constantly in a state of turnover and thus there is much matrix enzyme activity in health, disease, and tissue repair and remodelling. **It is evident that the activity of MMPs and their inhibitors is associated with tissue turnover as well as with gingivitis, destructive periodontitis and with the healing of the periodontal tissues following therapy.**

The periodontal ligament is one of the most metabolically active tissues in the body, and collagen metabolism represents most of this activity. The biological reason for this activity probably relates to its ability to adapt to occlusal forces generated during function. An important feature of connective tissues in general and the periodontal ligament in particular, is the process of constant renewal of the extracellular matrix components involving MMP. The regulation of extracellular matrix (ECM) turnover is influenced by the action of both TNF- α and TGF- β , in that TNF- α can promote the ECM degradation by enhancing expression of MMPs. This action is balanced by TGF- β which down regulates the secretion of these MMPs and enhances the production of their inhibitors (tissue inhibitor of matrix metalloproteases (TIMPs)). Normal homeostasis is maintained when MMPs are in equilibrium with inhibitors, the TIMPs. However, TIMPs can be degraded by P. gingivalis which could contribute to such dysregulation.

Cytokines

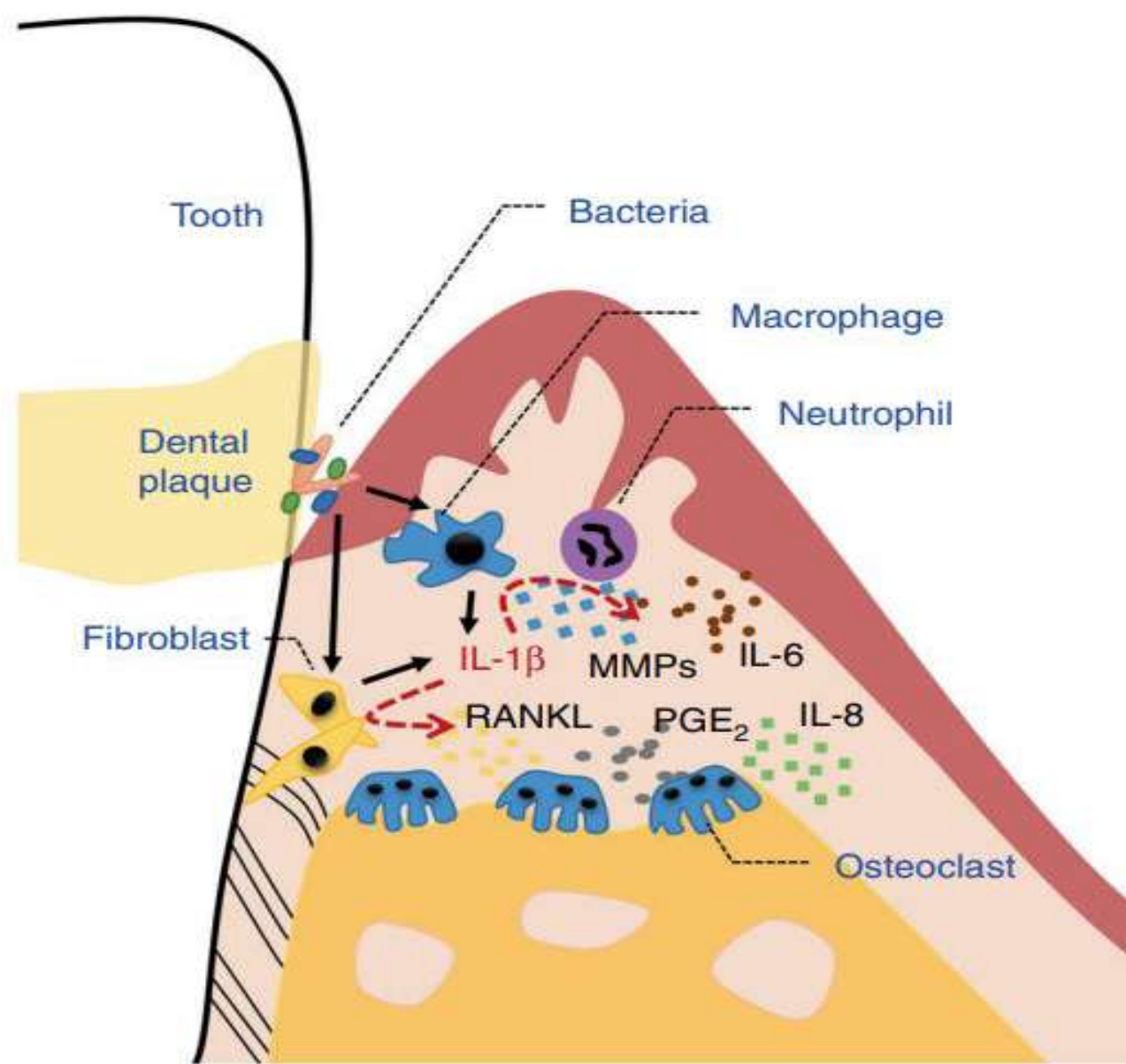
- Cytokines are soluble proteins, secreted by cells, which act as messenger molecules that transmit signals to other cells.
- They have numerous actions which include initiation and maintenance of immune and inflammatory responses and regulation of growth and differentiation of cells. The interleukins are important members of the cytokine group and are primarily involved in communication between leukocytes and other cells, such as epithelia, endothelia and fibroblasts, involved in both immune and inflammatory processes. These molecules are released in small amounts and have a variety of actions on cells which carry the specific receptor for the particular interleukin. Cytokines are numerous, many have overlapping functions and they are interlinked forming an active network which controls the host response.

Pro-inflammatory cytokines:

Cytokines such as interleukin (IL)-1 α , IL-1 β and tumour necrosis factor (TNF)- α stimulate bone resorption and inhibit bone formation.

Chemotactic cytokines:

A series of more than 20 molecules have been identified, among which the most famous and best characterized is **interleukin 8 (IL-8)**, which has powerful chemotactic functions for leukocytes particularly for neutrophils but also for lymphocytes and macrophages.

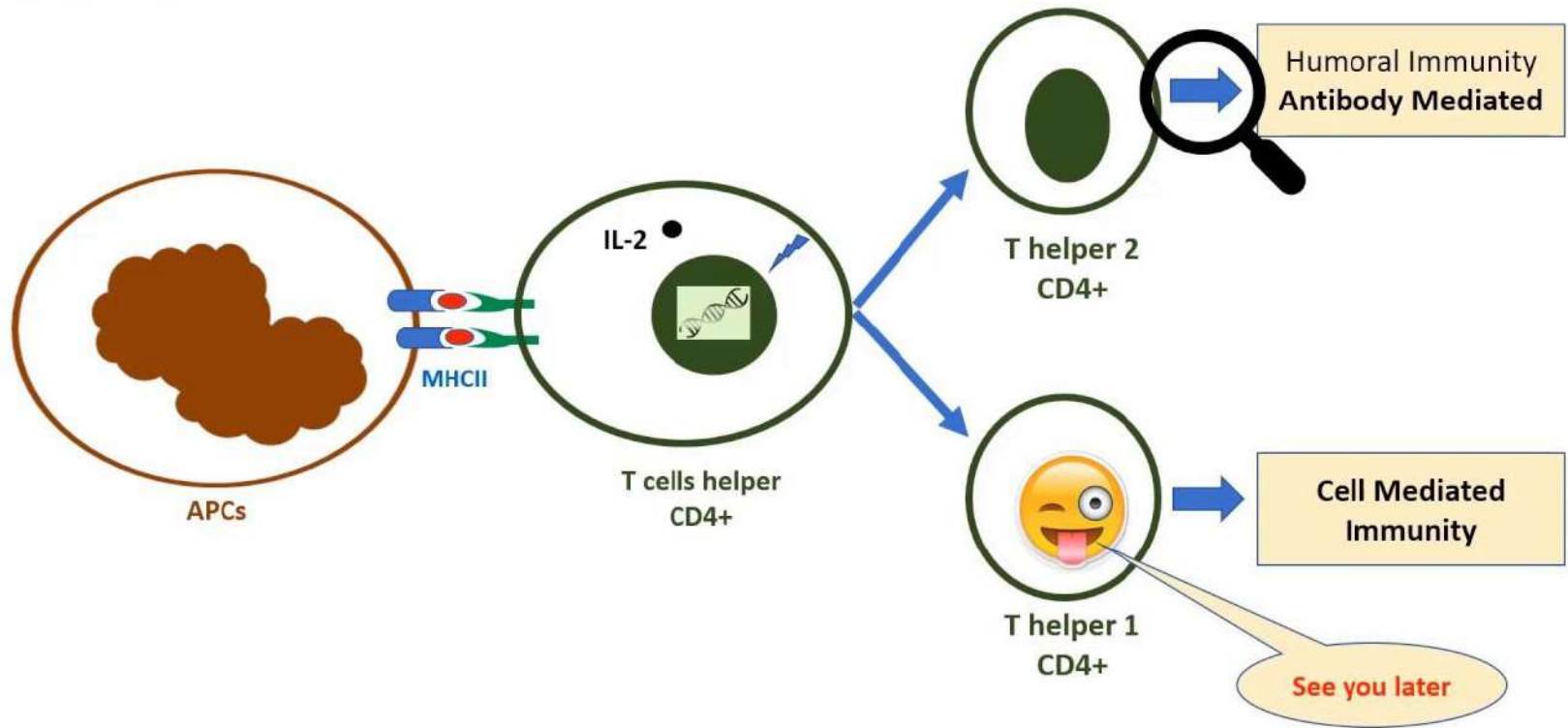


Role of IL-1 β in periodontitis. IL-1 β promotes the secretion of MMPs, RANKL, PGE₂, IL-6, IL-8, etc., which promotes osteoclastogenesis

Lymphocyte signaling cytokines:

T helper cells are lymphocytes within the tissues which regulate both the humoral and cell mediated immune responses via cytokines. The humoral immune response is promoted by a T helper cell type 2 (TH-2) which produces characteristic cytokines namely IL-5, IL-10 and IL-13. The TH-1 lymphocytes release IL-2 and interferon (IFN)- γ which enhance cell mediated responses. These cytokines provide a precise mechanism for the control of the immune response so that a sufficient response is produced to deal with the offending pathogen. Cytokines can influence the immune response through determining the class of immunoglobulin being produced, which may have quite a profound effect on antibody function.

● IL-2



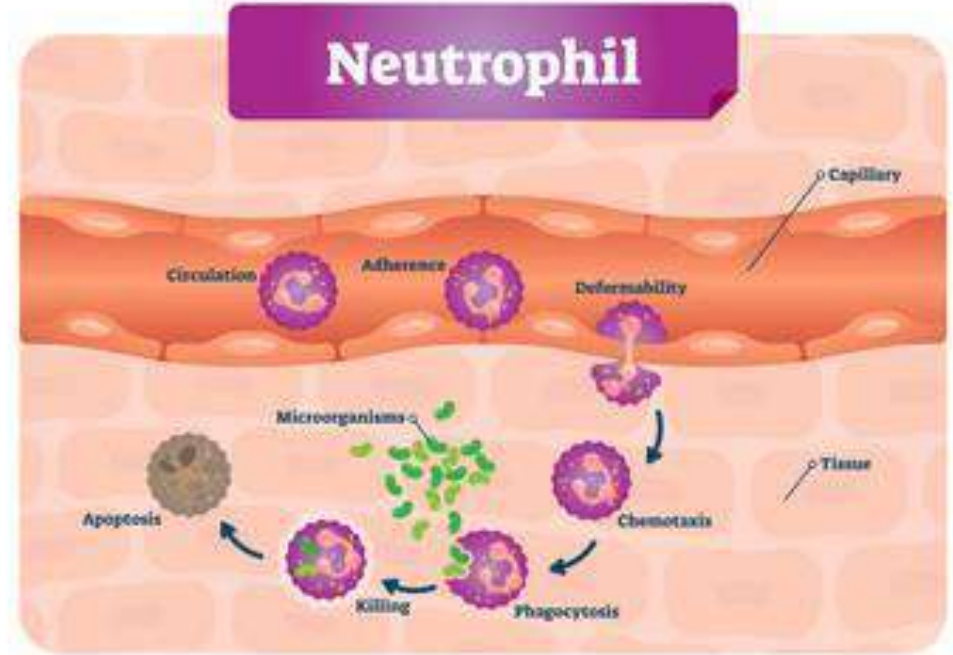
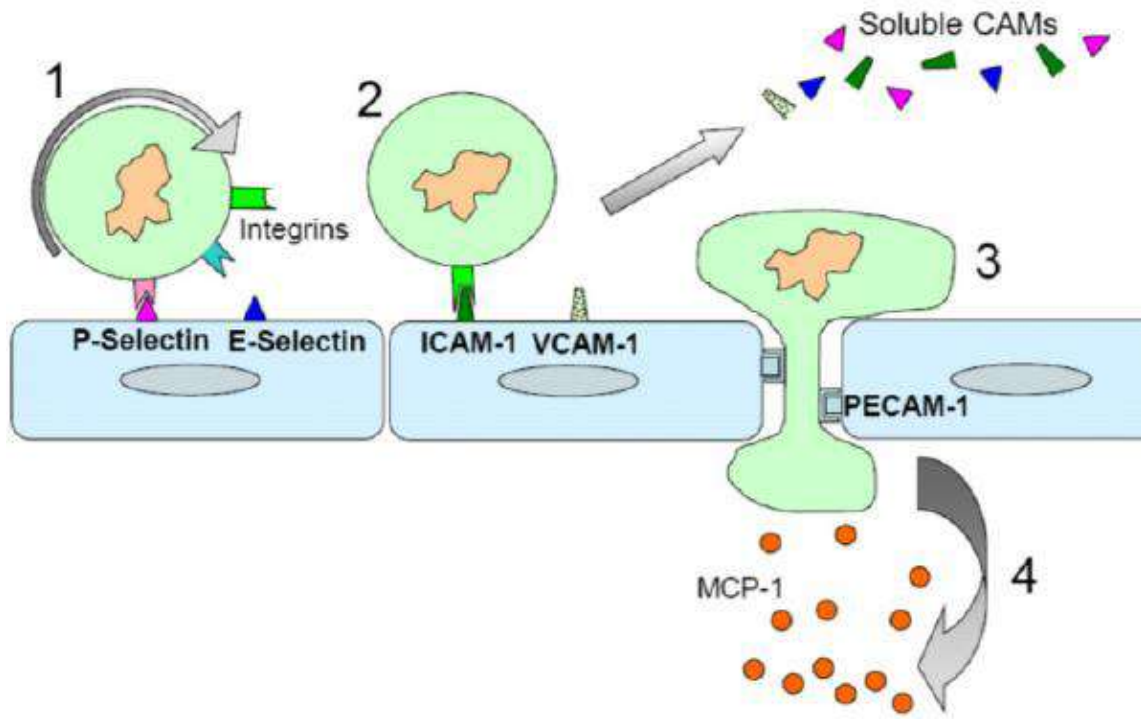
For example IgM molecules are more effective at bacteriolysis and IgG molecules are more effective at opsonization. The IgG antibodies exist as four distinct subclasses (IgG1, IgG2, IgG3 and IgG4) based on differences in the Fc portion of these molecules. The antibody subclass influences antibody function, IgG2 being less strong in binding antigen than IgG1. Several researchers have found IgG2 to be elevated over IgG1 in patients with severe periodontitis and propose that IgG subclass levels are important factors in susceptibility to periodontitis.

Prostaglandins

Prostaglandins are arachidonic acid derivatives which are important mediators of inflammation. The pro-inflammatory cytokines are capable of inducing macrophages and other cells to produce copious amounts of prostaglandins, particularly PGE2 which are **potent vasodilators** and **inducers of cytokine production** by various cells. **PGE2 acts on fibroblasts** and osteoclasts, together with cytokines, to **induce MMP production**, which is relevant to tissue turnover and in the **destructive process in periodontitis**. Many studies have examined the association of PGE2 with periodontal disease and suggest that its concentration in gingival crevicular fluid increases in gingivitis relative to health and is at very high concentrations during periods of periodontal disease progression.

Polymorph nuclear leukocytes (PMNs)

The PMN is the predominant leukocyte within the gingival crevice/pocket in both health and disease. PMNs from the circulation are attracted to the area via chemotactic stimuli elicited from microorganisms in the biofilm, and histologically PMNs can be seen traversing the gingival connective tissue in inflammation. Migration of leukocytes from the vessels into the gingival connective tissue, and through the junctional epithelium into the gingival crevice, is controlled via [adhesion molecules](#). Cellular migration involves three main structures: the endothelial cells, the cell adhesion molecules and the extravasating cells. Adhesion of leukocytes appears to be essential in controlling cellular traffic into inflamed areas and it has been proposed that cytokines may play an important role in regulation of this traffic



shutterstock.com · 1144289249

Host defense
processes

```
graph LR; A[Host defense processes] --> B[innate (non-specific)]; A --> C[adaptive (specific)]; B --> D[inflammatory response]; C --> E[immunological mechanisms]
```

The diagram is a flowchart with four teal rounded rectangular boxes. The first box on the left is titled 'Host defense processes'. Two lines branch from its right side to two boxes in the middle: 'innate (non-specific)' (top) and 'adaptive (specific)' (bottom). From the 'innate (non-specific)' box, a horizontal line connects to a box on the right titled 'inflammatory response'. From the 'adaptive (specific)' box, a horizontal line connects to a box on the right titled 'immunological mechanisms'.

innate (non-specific)

inflammatory
response

adaptive (specific)

immunological
mechanisms

THE INNATE DEFENSE SYSTEMS

Innate immune mechanisms operate without any previous contact with the disease causing microorganism. These mechanisms include **physical barriers** of the oral mucosal epithelial surfaces, vascular and cellular aspects of the **inflammatory responses**. The epithelial surface is the first region of the periodontium which comes into contact with and responds to bacteria attaching and colonizing the dento- gingival region. Prevention of attachment and colonization is important for the host defenses. The oral mucosa itself is not simply a barrier but has a chemical composition which may be detrimental to bacteria. Furthermore, the cells of the epithelium can respond to the bacteria by (1) producing and/or releasing cytokines and other molecules that kill the microbes and (2) releasing other molecules (such as IL-1) capable of inducing or enhancing the inflammatory reaction. The epithelium can also respond by increasing expression of surface molecules such as cell adhesion molecules which can function with cytokines and chemo attractants to bring leukocytes to the region.

The major functions of the innate immune system include:

- ❑ Acting as a **physical and chemical barrier** to infectious agents.
- ❑ **Recruiting immune cells to sites of infection**, through the production of chemical factors, including specialized chemical mediators, called cytokines
- ❑ **Activation of the adaptive immune system** through a process known as antigen presentation.
- ❑ **Activation of the complement cascade** to identify bacteria, activate cells, and promote clearance of antibody complexes or dead cells.
- ❑ **Identification and removal of foreign substances** present in organs, tissues, the blood and lymph, by specialized white blood cell.

Adaptive immune system

The adaptive immune system, also known as the acquired immune system or, as the specific immune system, is a subsystem of the overall immune system that is composed of highly specialized, systemic cells and processes that eliminate or prevent pathogen growth.

Adaptive immunity creates immunological memory after an initial response to a specific pathogen, and leads to an enhanced response to subsequent encounters with that pathogen.

Unlike the innate immune system, the adaptive immune system is highly specific to a particular pathogen.

Functions

1. Recognition of specific antigens during the process of **antigen presentation**.
2. Generation of responses that are tailored to maximally eliminate specific pathogens or pathogen-infected cells.
3. Development of immunological memory, in which pathogens are "remembered" through memory B cells and memory T cell

Thank you

est of gum

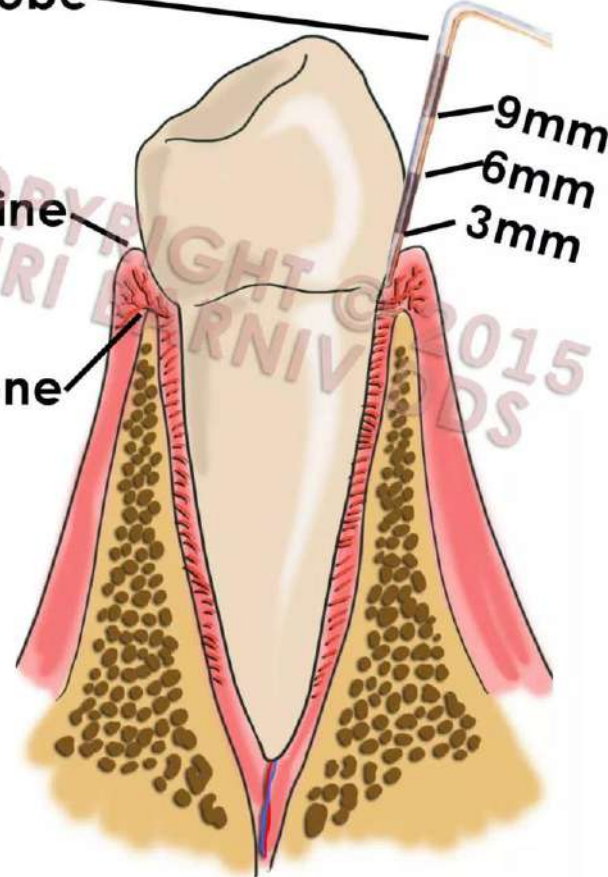
st of jaw

NORMAL HEALTHY GUMS AND BONE

Periodontal probe

Crest of gum line

Crest of jaw bone



9mm
6mm
3mm

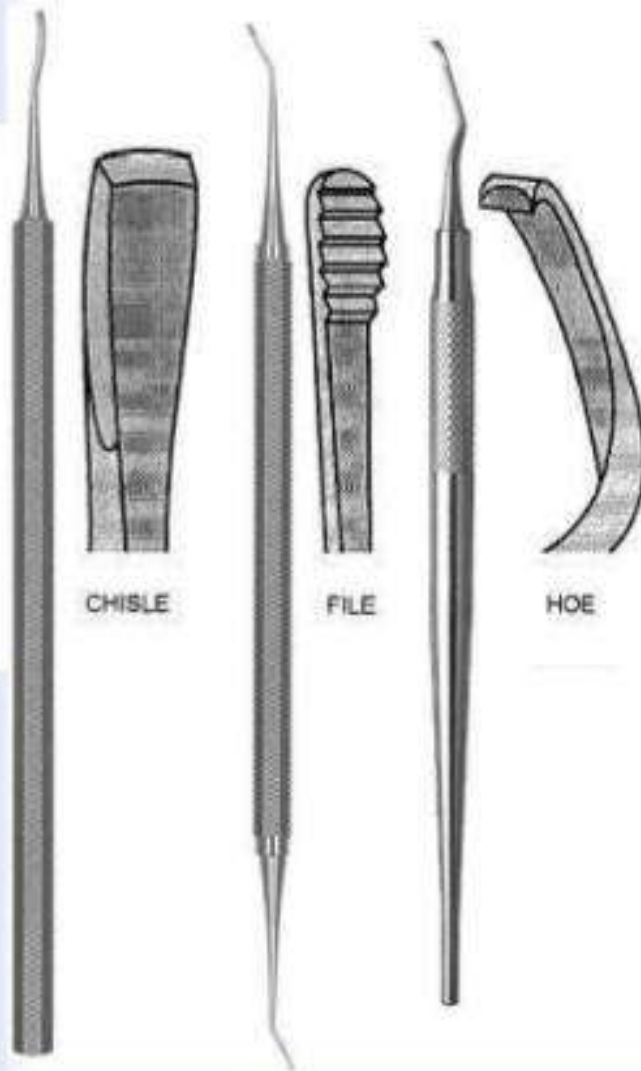








SUBGINGIVAL SCALERS



Chisel scaler

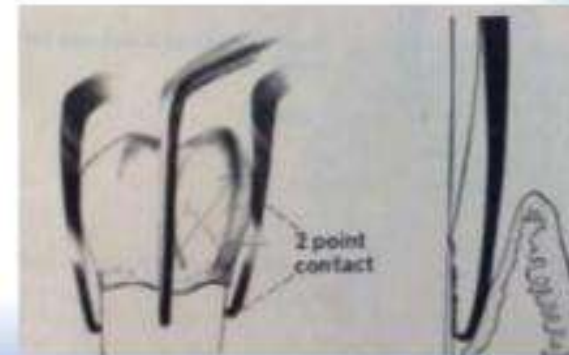
Used in interproximal area
Used in a push motion

File

Used to crush large pieces of
calculus deposits.

Hoe

Efficient in removing subgingival calculus
Blade is beveled at 45 degrees
Working end is bent at an angle of 99
degrees to shank.

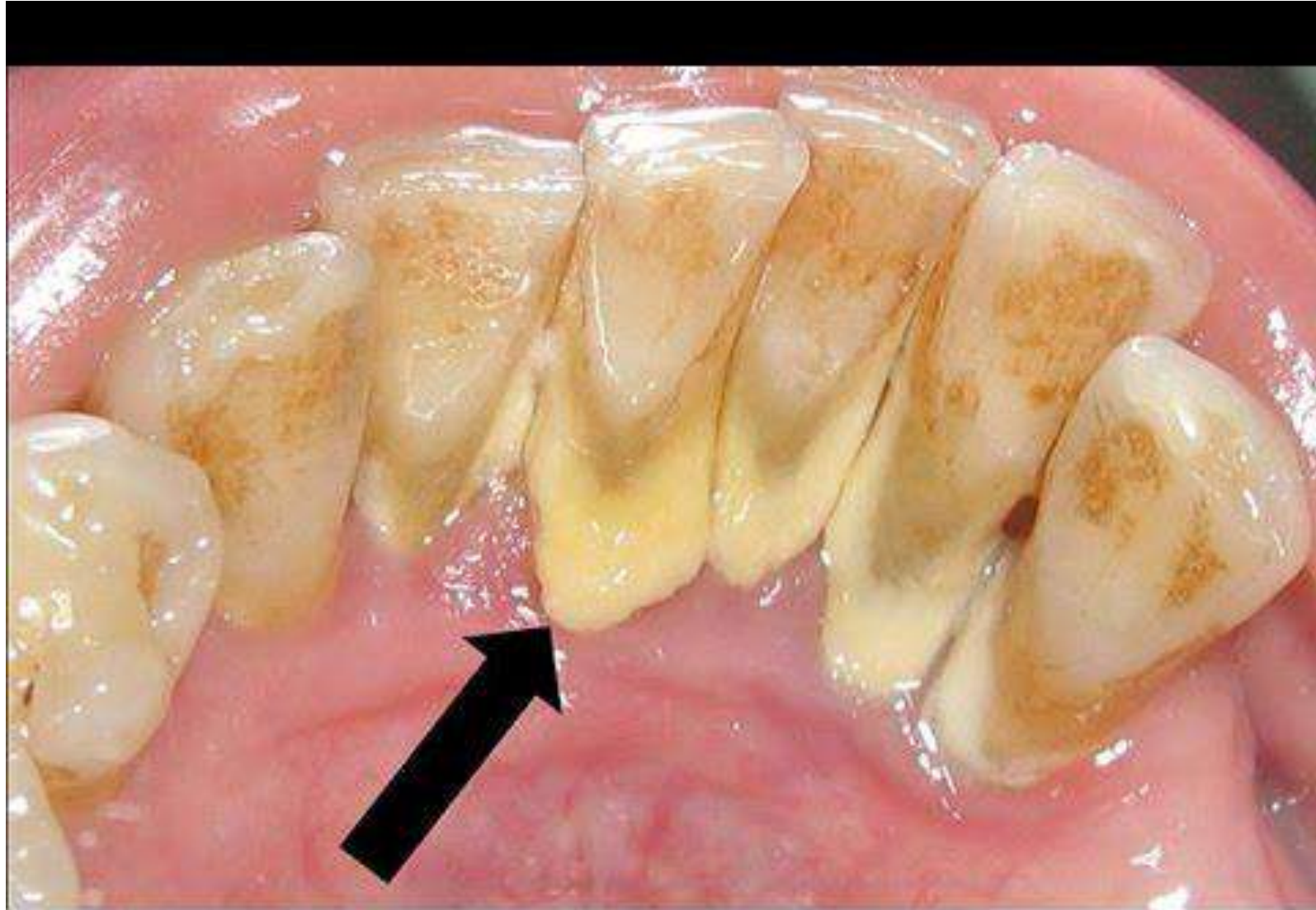






eshopbid666





SCALING, ROOT-PLANING AND CURETTAGE INSTRUMENTS

A. Curette : for sub gingival scaling, root planing and removal of the soft tissue lining the pocket

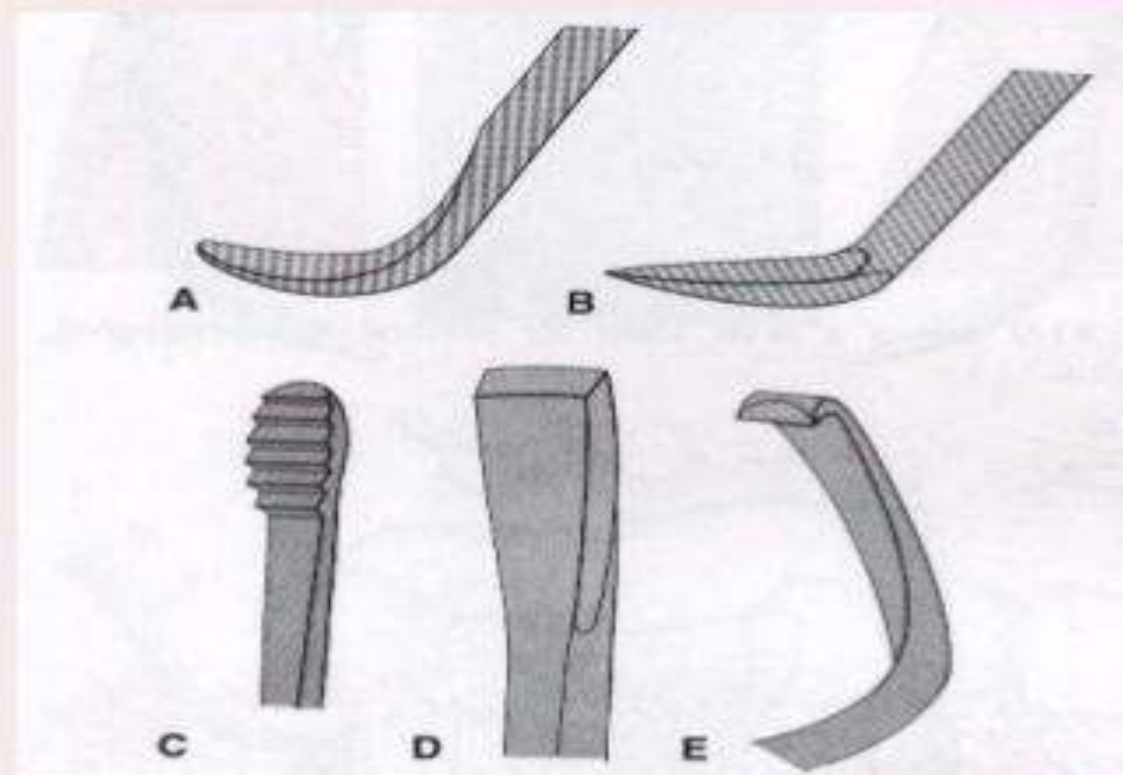
B. Sickle Scaler : to remove supra gingival calculus

C. File

D. Chisel

E. Hoe

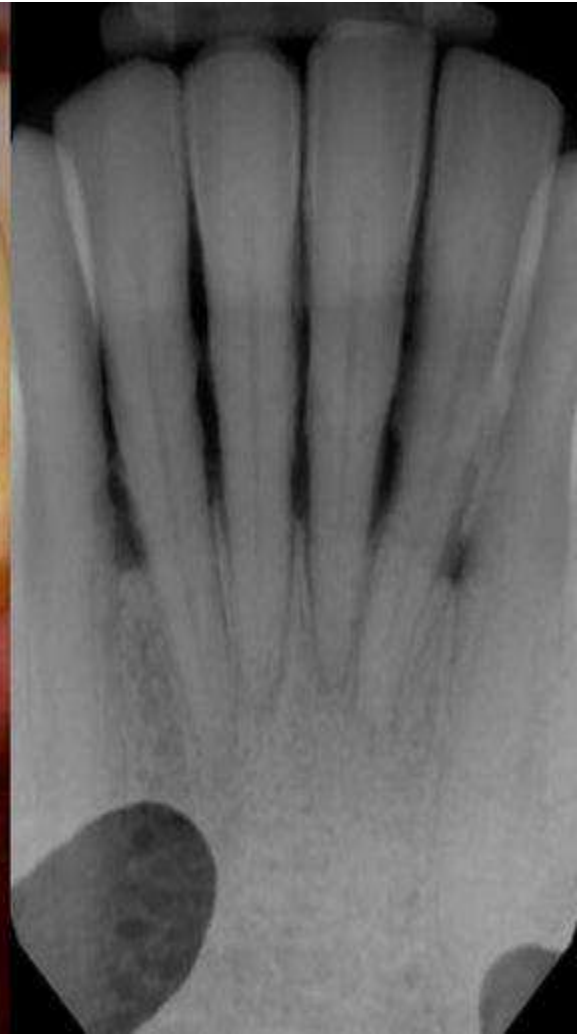
to remove
tenacious
calculus and
altered
cementum





BEFORE

AFTER





Procedure: SCALING ROOT PLANING









PUMICE POWDER

Grain size: 120mesh (5rO48)

Mohs hardness: 4.8

PH: 7

Oil No: 1302-05-B











Snap-on



Screw type



Latch type

dentp.en.alibaba.com



Tapered

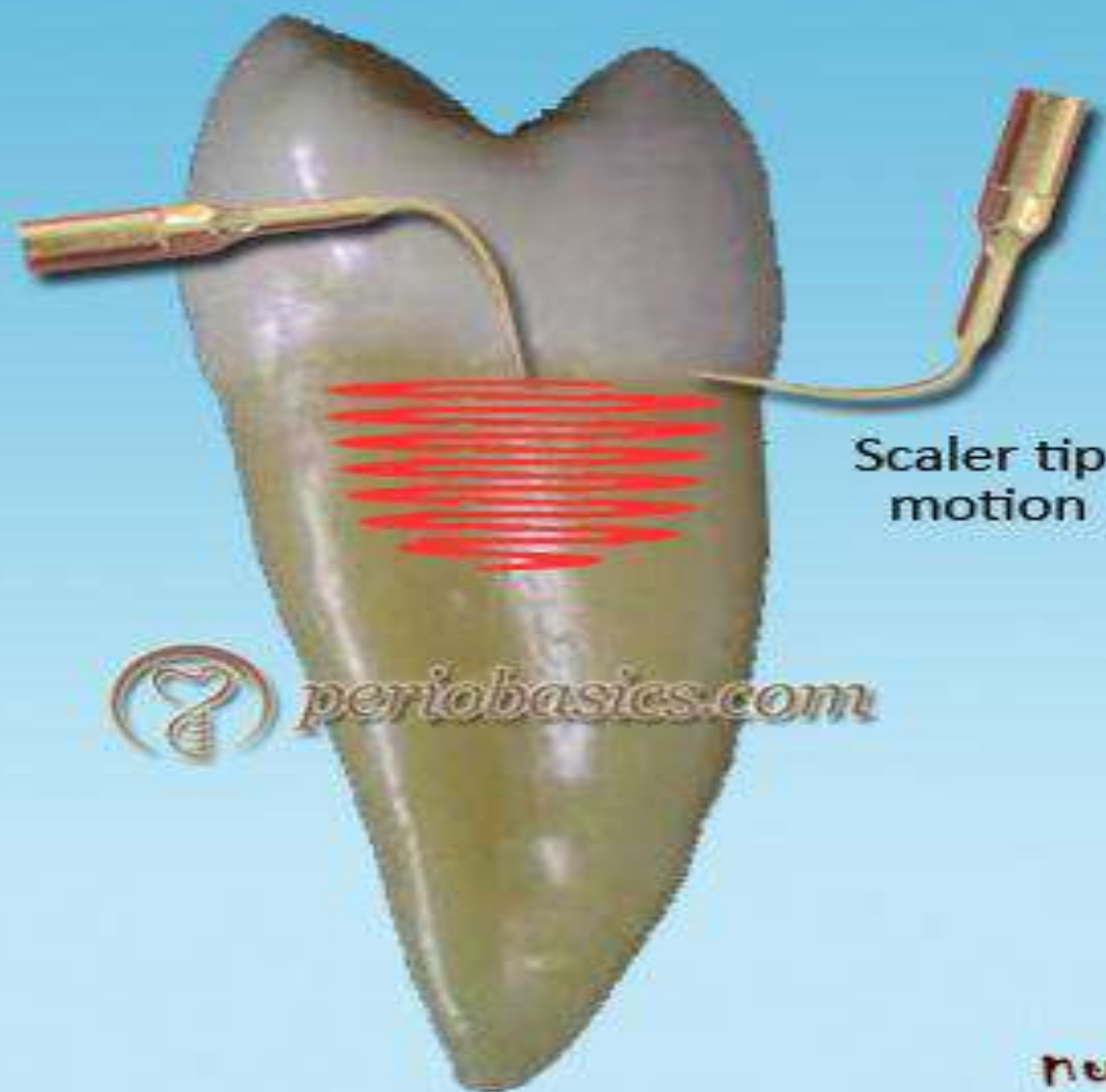


Flat



Cup-shape



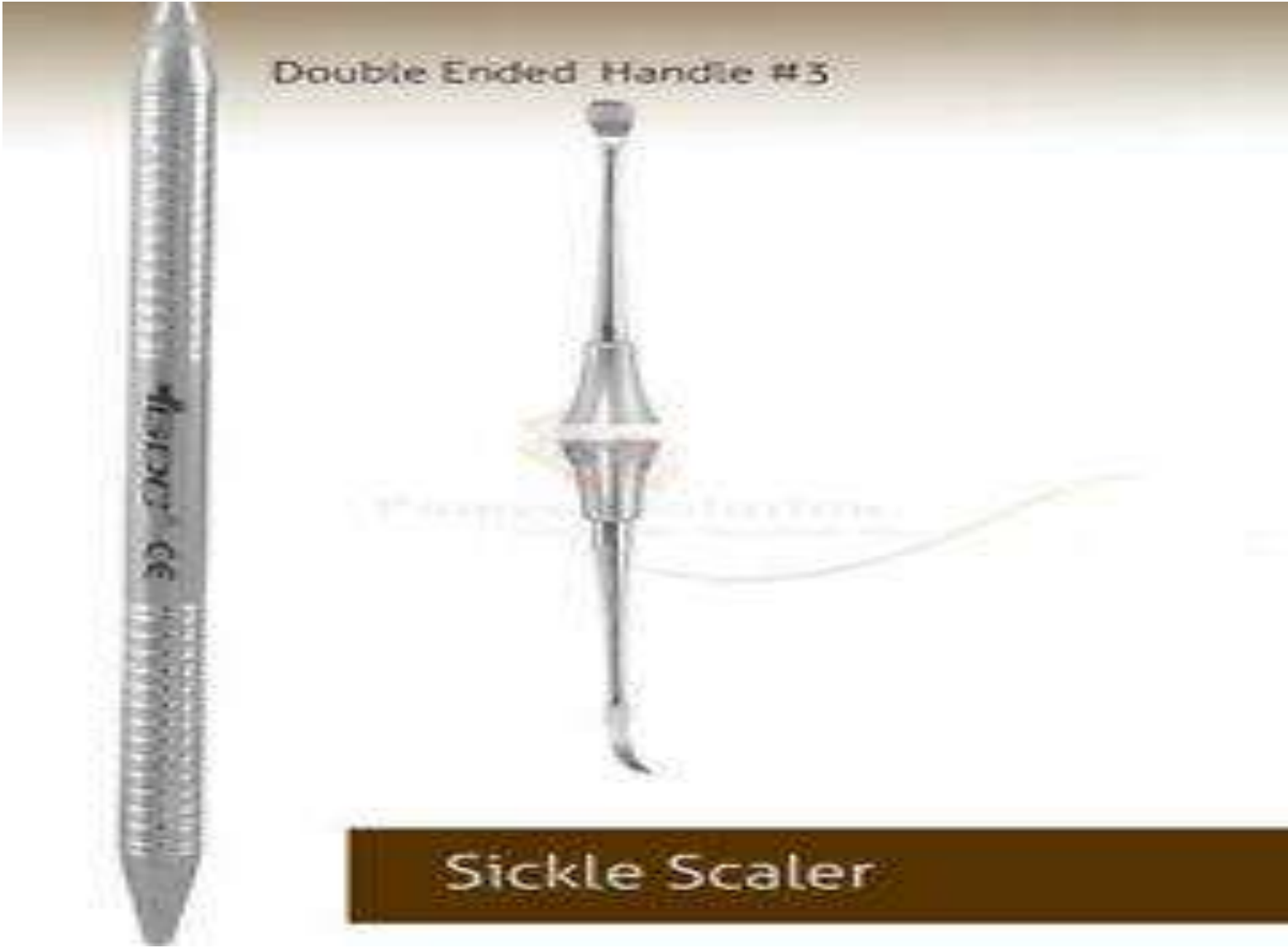


Scaler tip
motion

 periobasics.com

nue49









AR INSTRUMENTED

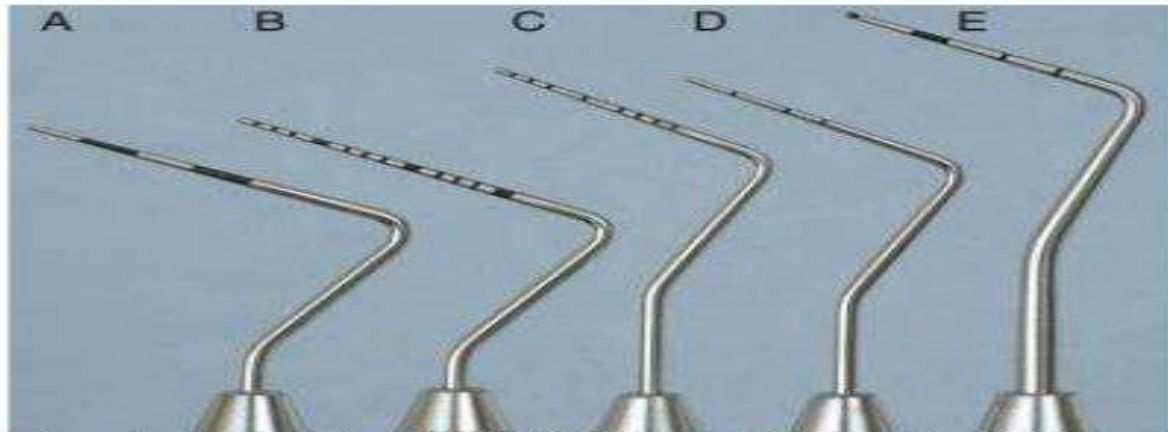
Periodontics Instruments

Explorers

- Use: **1. Locate Sub-gingival Deposits and Caries.**
2. Check Smoothness of root surface after planning.
3. Assess Restorative problems.
- Thin, Flexible, Wire-like working end. Taper to sharp point.
- Curved, Right-angled & Area specific.

Periodontal Probes

- Use: **1. Locate & Measure depth of pocket and determine it's configuration.**
2. Assess Loss of Attachment.
3. Detect Sub-gingival Deposits.
- Tapered with blunt round tip, mm markings for accuracy.
- Ball-end to avoid penetration into junctional epithelium.
- Diameter less than or equal 0.6 mm.
- Probing Force more than 0.25 N traumatize healthy tissue. (25-50g for clinic)
- Ball-end to avoid penetration into junctional epithelium.



A, Marquis color-coded probe. **B**, UNC-15 probe. **C**, University of Michigan "O" probe, with Williams markings. **D**, Michigan "O" probe with markings at 3, 6, and 8 mm. **E**, World Health Organization (WHO) probe.





Hoe scaler





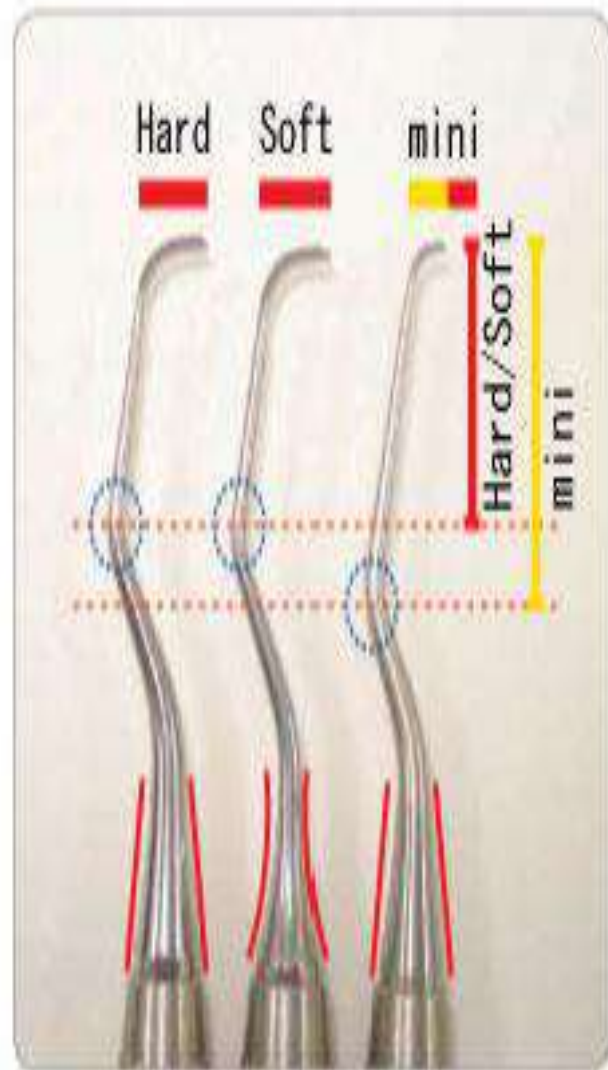
1.4mm



HSA12-13

• Anterior Hoe Scaler

For anterior buccal and lingual surfaces.







Microbiologic Specificity of Periodontal Diseases

م. سها أسود دهش العزاوي

B.D.S, MSc. Periodontology

- 1. Traditional Nonspecific Plaque Hypothesis.**
- 2. Specific Plaque Hypothesis.**
- 3. Updated Nonspecific Plaque Hypothesis.**
- 4. Ecologic Plaque Hypothesis.**
- 5. Keystone Pathogen Hypothesis.**

Traditional Nonspecific Plaque Hypothesis

- Periodontal noxious products by the entire plaque flora are responsible in a proportional way to the severity of the gingival inflammation.
- According to this thinking, when only small amounts of plaque are present, the noxious products are neutralized by the host. Similarly, large amounts of plaque would produce large amounts of noxious products, which would essentially overwhelm the host's defenses. The NSPH have focused the **quantity of plaque** that determined the pathogenicity without discriminating between the levels of virulence of bacteria

- Believing this, the host would have a **threshold capacity** to detoxify bacterial products (e.g., saliva neutralizing acid) and disease would only develop if this threshold was surpassed and the virulence factors could no longer be neutralized.
- **The conclusion** was that if any plaque has an equal potential to cause disease, the best way of disease prevention would be non-specific mechanical removal of as much plaque as possible by e.g., tooth brushing

- **Several** observations contradicted these conclusions. **First**, some individuals with considerable amounts of plaque and calculus, as well as gingivitis, never developed destructive periodontitis. **Furthermore**, individuals who did present with periodontitis demonstrated considerable site specificity in the pattern of disease. Some sites were unaffected, whereas advanced disease was found in adjacent sites. In the presence of a uniform host response, these findings were inconsistent with the concept that all plaque was equally pathogenic. Recognition of the differences in plaque at sites of different clinical status (i.e., disease versus health) led to a renewed search for specific pathogens in periodontal diseases and a conceptual transition from the nonspecific to the specific plaque hypothesis.

- **In addition**, the improvement of techniques to isolate and identify bacteria in the mid-20th century led to the abandoning of the NSPH. Although the nonspecific plaque hypothesis has been discarded in favor of the specific plaque hypothesis or the ecologic plaque hypothesis, much clinical treatment is still based on the nonspecific plaque hypothesis through mechanical plaque removal that represents the most efficient way of preventing disease.

Specific Plaque Hypothesis

- The specific plaque hypothesis states that only certain plaque is pathogenic, and its pathogenicity depends on the presence of or increase in specific microorganisms. This concept predicts that plaque harboring specific bacterial pathogens results in a periodontal disease because these organisms produce substances that mediate the destruction of host tissues. Acceptance of the specific plaque hypothesis was spurred by the recognition of *A. actinomycetemcomitans* as a pathogen in localized aggressive periodontitis.

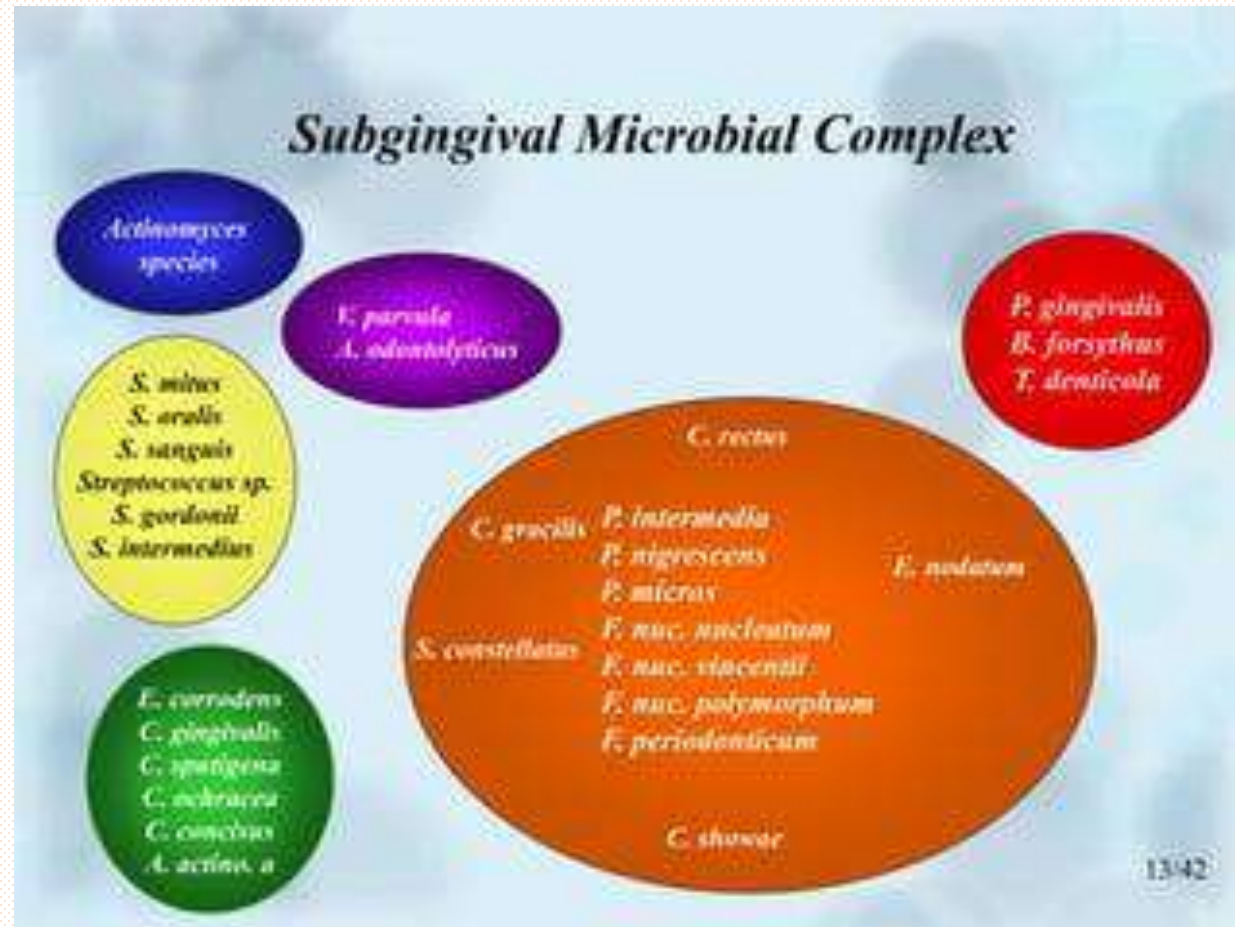
Specific Plaque Hypothesis

- In the 1970s, culture-based techniques and microscopy allowed discrimination of specific bacterial species and opened the hunt for disease-related micro-organisms.
- It was noticed that the antibiotic kanamycin was particularly effective against cariogenic species such as oral streptococci and reduced caries formation. This suggested that removing cariogenic bacteria from the oral cavity using antibiotics could prevent caries. In 1976, Walter J. Loesche announced the “Specific Plaque Hypothesis” (SPH), postulating that dental caries was an infection with specific bacteria in the dental plaque of which the most relevant were “mutans streptococci” (main species: *Streptococcus mutans* and *Streptococcus sobrinus*) and *Lactobacilli*.
- This hypothesis proposed that use of antibiotics against specific bacterial species could cure and prevent caries.

- However, results from clinical studies, then and today, are not very promising. For instance, even though the use of kanamycin resulted in an overall reduction of caries, at some surfaces the caries rate increased. This indicates that kanamycin failed to penetrate certain niches allowing cariogenic species to have a selective advantage and accumulate there. Furthermore antibiotics reduced the abundance of cariogenic bacteria but failed to eliminate them thus as soon as the treatment was stopped, abundance increased, while a long period of treatment leads to antibiotic resistance.

- The development of the anaerobic hood in the 1970s for the first time allowed cultivation of the strict anaerobic species. This extended the SPH to periodontal diseases which were proposed to be inflammations caused by specific periopathogens and antibiotic treatment would be effective. However, in line with the use of antibiotics in caries treatment,
- Recent clinical studies evaluating the effectiveness of antibiotics as adjunct in periodontal therapy have not booked significant success either.
- **Potential periopathogens included:** spirochetes, streptococci, and actinomyces. In addition, Gram-negative, anaerobic rods including black-pigmented *Bacteriodes* such as *Prevotella melaninogenica* and *Campylobacter* and facultative anaerobic, Gram-negative rods of the genera *Capnocytophaga*, *Eikenella* and *Actinobacillus* were identified as periopathogens. However, these findings were limited due to the large number of uncultivable species (~50%) and the bias toward easily cultivable species.

- The finding of different species related to periodontal disease led to the idea that oral disease could be initiated by a number of specific pathogens. This idea was further investigated over the next decades and led to the famous **Socransky-complexes** which include bacterial clusters based on their association with periodontal disease.



Updated Nonspecific Plaque Hypothesis

- Theilade also noticed that the “specific-pathogens” from the SPH were indigenous bacteria and sometimes common bacteria in health, which led to an updated NSPH in 1986 focusing on periodontal disease. At this time most researchers seemed to agree that gingivitis was a non-specific inflammatory reaction to a complex indigenous microbiota. However, the updated NSPH took into consideration that some indigenous subgingival bacteria can be more virulent than others and that plaque composition changes from health to disease.

- Nevertheless, it stated that all bacteria in plaque contribute to the virulence of the microflora by having a role in either colonization, evasion of the defense mechanism, and/or provocation of inflammation and tissue destruction. Theilade's statement that "any microbial colonization of sufficient quantity in the gingival crevice causes at least gingivitis" was supported by the fact that a non-pathogenic plaque (i.e., not causing gingivitis in the absence of oral hygiene) had never been observed.
- Additionally, it was considered that some people have gingivitis for a lifetime without tissue and bone destruction, while others encounter rapid progression into periodontitis. Unlike the classic NSPH, the updated NSPH could explain this by taking into account that differences in the plaque microbial composition could lead to differences in pathogenic potential.

Ecologic Plaque Hypothesis

- In 1994 Philip D. Marsh proposed a hypothesis that combined key concepts of the earlier hypotheses. In his “Ecological Plaque Hypothesis” (EPH), **disease is the result of an imbalance in the total microflora due to ecological stress, resulting in an enrichment of some “oral pathogens”** or disease-related micro-organisms. This idea was not entirely new since Theilade , in the review proposing the U-NSPH concluded that “increased virulence of plaque (leading to disease) is due to a plaque ecology unfavorable to the host and favorable for overgrowth by some of the indigenous bacteria having a pathogenic potential”.
- Importantly, **Marsh expanded this theory** and related the **changes in microbial composition to changes in ecological factors such as the presence of nutrients and essential cofactors, pH and redox potential** (Marsh, 1994, 2003).

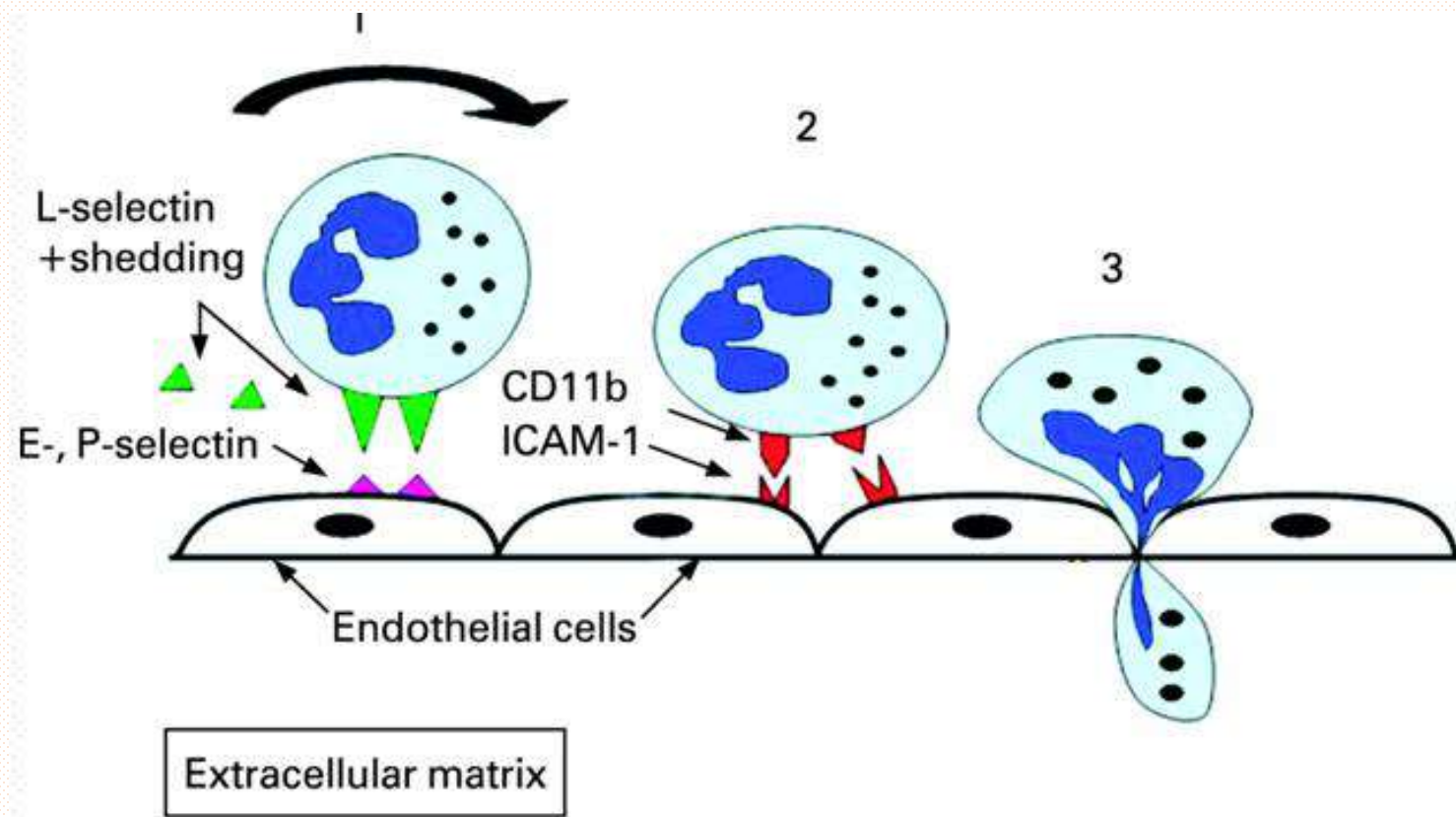
- For example, frequent exposure to a low pH, for instance as the result of sugar fermentation, leads to a relative increase of acid-tolerant species. The thought arose that disease could be prevented by interfering with processes that break down homeostasis and change composition. For example, non-fermentable sweeteners could be used to replace sugar and thus prevent acidification.
- Thus, the classical “everything is everywhere, but, the environment selects” was successfully applied to dentistry.
- Marsh also considered the reverse: **the bacteria in dental plaque affect the environment**. For instance, early colonizers of supragingival dental surfaces, are usually facultative anaerobic bacteria that use the oxygen, producing carbon dioxide and hydrogen. This lowers the redox potential giving strict anaerobes a chance to settle and multiply in the biofilm.

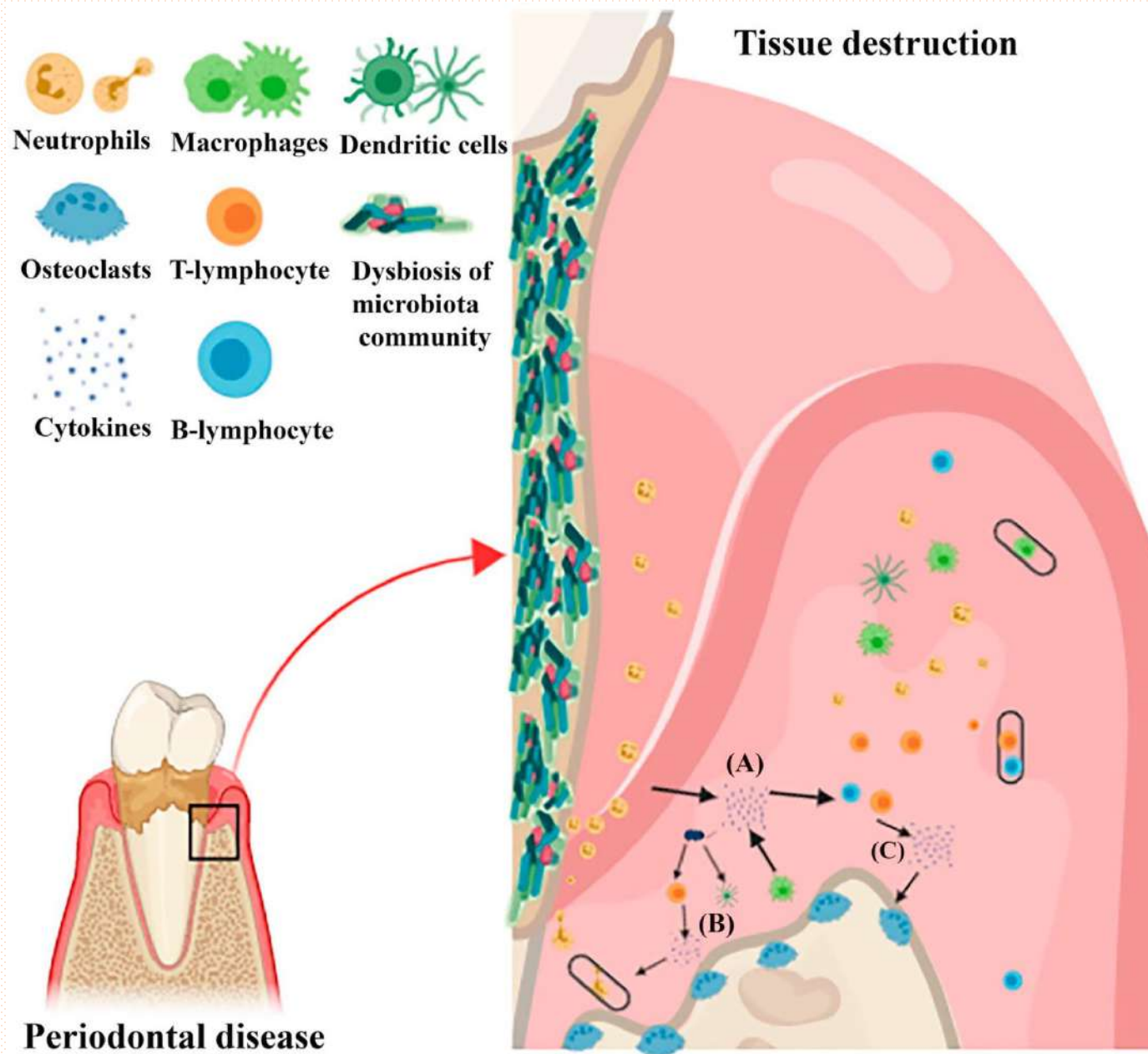
Keystone Pathogen Hypothesis

- Certain species have an effect on their environment that is disproportional relative to their overall abundance.
- The Keystone-Pathogen Hypothesis” (KPH) indicates that **certain low-abundance microbial pathogens can cause inflammatory disease by increasing the quantity of the normal microbiota and by changing its composition.** For instance, *Porphyromonas gingivalis* is shown to be able to manipulate the native immune system of the host. By doing so it was hypothesized that it does not only facilitate its own survival and multiplication, but of the entire microbial community.

- In contrast to dominant species that can influence inflammation by their abundant presence, keystone pathogens can trigger inflammation when they are present in low numbers. When disease develops and advanced stages are reached, the keystone pathogen are detected in higher numbers. Importantly, even though their absolute number increases, keystone pathogens can decrease in levels compared to the total bacterial load which increases as plaque accumulates in periodontitis.
- The KPH was developed by observing the properties of the “red complex” bacterium *P. gingivalis*.
- Studies in mouse models showed that very low presence of *P. gingivalis* (<0.01% of the total bacterial count in plaque) could alter the plaque composition, leading to periodontitis. In germ-free mice, *P. gingivalis* was able to colonize by itself, but was not able to trigger disease without the presence of other bacterial species. This indicates that (some of) the commensal microbiota is essential in the disease process. Evidence of *P. gingivalis* acting as a key stone pathogen was also obtained in rabbit models and non-human primates.

- The role of the host-immune system is critical in the KPH. At health, periodontal tissue contains a **wall of neutrophils**, between the plaque and the epithelial surface, residing just outside the epithelial cells. Expression of mediators such as **interleukin 8 (IL-8)**, **intercellular adhesion molecule (ICAM)** and **E-selectin** is required to form this neutrophil wall. E-selectin is required for neutrophil migration from the highly vascularized gingival tissue, IL-8 is a key neutrophil chemo-attractant produced by epithelial cells, and ICAM facilitates adhesion of neutrophils to the tissue allowing formation of this wall. Furthermore, the epithelium expresses low levels of a wide range of toll-like receptors (TLR's), including TLR1-TLR9 that mediate the response to a broad range of microorganisms.
- The array of different TLRs in combination with the multitude of bacterial species lead to a large variety of cytokines that are expressed at health.





- Studies in germ-free mice show that there are low levels of innate host mediators, such as IL-1B, present in the periodontal tissue. This indicates that a basic level of cytokine expression is genetically programmed without bacterial challenge. **The composition and amount of bacteria in plaque modifies cytokine expression further.**

- Evidence was found of three major KPH mechanisms of *P. gingivalis* that could impair the above mentioned host defenses:
- (1) Toll-like receptor (TLR) response manipulation,
- (2) interleukin 8 (IL-8) subversion and
- (3) the corruption of the complement system.

- *In vitro*, the TLR response is manipulated by *P. gingivalis* with the help of two types of lipopolysaccharides (LPS) with different lipidA structures Pg LPS (type I) and Pg LPS (type II). Type I is a TLR4 agonist thus activating the immune system, while Type II is a TLR4 antagonist inhibiting the immune response to *P. gingivalis*. The concentration of iron determines which type of LPS is expressed. In the oral cavity, the main source of iron is heme, found in the gingival crevicular fluid (GCF). During inflammatory process, GCF increases stimulating *P. gingivalis* type II LPS expression, thus reduces the TLR4 response. It was proposed that this could facilitate survival and multiplication of the entire microbial community.

- *Porphyromonas gingivalis* can block production of IL-8 *in vitro*, which is produced by gingival epithelial cells in response to other bacteria, by secreting a **serine phosphatase** that **inhibits the synthesis of IL-8**. This process is called “**local chemokine paralysis**” and delays the recruitment of neutrophils preventing proper neutrophil wall formation, of which was proposed that it could facilitate initial microbial colonization of the periodontium. Other “red complex” bacteria such as *T. denticola*, are also able to manipulate the IL-8 response of the host.

- The third and best *in vivo* documented key stone pathogen mechanism is the interference with the complement system.
- To be a successful pathogen in humans (and any other mammal) a microorganism needs to be able to avoid complement-mediated detection and killing. Again, the best-studied example in the oral cavity is *P. gingivalis* that produces **membrane bound and soluble arginine-specific cysteine proteinases called “gingipains”**. Gingipains can cleave complement factors C3 and C5 into active fragments C5a (cell activator) and C3b (phagocytosis enhancer). These fragments can be further degraded by gingipains resulting in loss of their function. More relevant is that in the presence of gingipains the levels of the inflammatory mediator C5a increase within seconds. This leads to an increased activation of the C5a receptor (C5aR) on leukocytes.

- C5aR is involved in cross talk with TLR2, which is activated in parallel by *P. gingivalis* (and other bacterial) surface ligands. While this crosstalk leads to increased inflammation, it impairs the killing capacity for leukocytes. In mouse models this mechanism has a major role in accelerating periodontitis development and bone loss. A *P. gingivalis* strain that lacks gingipains failed to change the oral microbiota and induce bone loss. Additionally, periodontitis did not develop in mice lacking one of the two involved receptors C5aR or TLR2. This provides clear evidence that in mice the dysbiosis caused by *P. gingivalis* is mainly due to complement subversion.

- In conclusion, it was proposed that currently known and unknown keystone pathogens use a combination of these and presently unknown mechanisms to manipulate the innate defense system leading to destructive periodontitis.

Thank you

Pathogenesis of periodontal disease


م. سها أسود دهش العزاوي

B.D.S, MSc. Periodontology

- Inflammatory and immune reactions to microbial plaque are the predominant features of gingivitis and periodontitis.
- Inflammatory and immune processes operate in the gingival tissues to protect against local microbial attack and prevent microorganisms or their damaging products from spreading into or invading the tissues. These host defense reactions are, however, also considered potentially harmful to the host in that inflammation can damage surrounding cells and connective tissue structures.
- These "defensive" processes could therefore paradoxically contribute to the tissue injury observed in gingivitis and periodontitis.

The inflammatory reaction is visible both **clinically** and **microscopically** in the affected periodontium.

INITIATION OF PERIODONTAL DISEASE

- **Pathogenesis**  biological and histological events that occur in the tissue during the process of conversion from the healthy state to diseased one.
- Most normal subjects maintaining a high standard of oral hygiene are not likely to develop advanced periodontal disease.
- Microorganisms quickly start to colonize clean tooth surfaces once an individual abstains from mechanical tooth cleaning; within a few days microscopical and clinical signs of gingivitis are then apparent. These inflammatory alterations are resolved or reversed when adequate tooth cleaning measures are resumed. The inflammatory changes may remain confined to the gingival area for several years, but at some sites gingivitis eventually shifts to destructive periodontal disease resulting in loss of connective tissue attachment and alveolar bone.
- Clearly some imbalance of the host-microbial relationship is occurring in the destructive lesions, which may be unique to that site and to periodontally susceptible individuals.

MECHANISMS OF PATHOGENICITY

For a periodontal pathogen to cause disease, it is essential that the pathogen be able to

(1) Colonize the subgingival area and

(2) Produce factors that either directly damage the host tissue or lead to the host tissue damaging itself.

To colonize subgingival sites, a species must be able to:

(1) Attach to one or more of the available surfaces,

(2) Multiply,

(3) Compete successfully against other species, and

(4) Defend itself from host defense mechanisms.

Adhesions

To establish in a periodontal site, a species must be able to attach to one or more surfaces including the **tooth**, the **sulcular or pocket epithelium** or other **bacterial species attached to these surfaces**. Some of the **adhesins** that have been identified on subgingival species include fimbriae and cell associated proteins. **Receptors** on tissue surfaces that species adhere to them include galactosyl residues and proline rich proteins.

Coaggregation

While many species attach directly to host surfaces, other species attach to bacteria attached to such surfaces. This phenomenon is called coaggregation.

Multiplication

The gingival crevice and/or periodontal pocket might be considered a lush area for microbial growth, but is in fact a rather stringent environment for a bacterial species to live. The mean temperature of the area averages about 35°C and ranges from 30'-38"C. The pH is rather restricted (pH 7. 0-8.5).

Three sources of nutrient are available to subgingival organisms (diet, host and other subgingival species). Certain nutrients essential to some bacterial species must be formed by other species in that area. However, the precursors to such substances and certain specific growth factors such as hemin must be derived from the host. Gingival crevice fluid is not particularly rich in nutrients, creating a major competition for the small amounts available. In addition, nutrients delivered in relative abundance to the outer layers of plaque may not reach deeper layers.

Interbacterial relationships

Bacterial interactions play important roles in species survival. Some inter-species relationships are **favorable**, in that one species provide growth factors or facilitate attachment of another. Other relationships are **antagonistic** due to competition for nutrients and binding sites or to the production of substances which limit or prevent growth of a second species such as production of hydrogen peroxide by *S. sanguis* which suppress the growth of *A. actinomycetemcomitans*. On the other hand, the growth of *S. sanguis* has been shown inhibited by a bacteriocin produced by *A. actinomycetemcomitans*.

Overcoming host defence mechanisms

Subgingival plaque microorganisms appear to overgrow and lead to severe disease in immune-compromised hosts, particularly those with neutrophil disorders. A bacterial species has a number of host-derived obstacles to overcome when colonizing a subgingival site' These include the **flow of saliva** and **gingival crevice fluid** and **mechanical displacement** by chewing and speaking. **Substances in saliva and gingival crevice fluid** may aid in the prevention of colonization by blocking the binding of bacterial cells to mammalian surfaces' Such factors include specific antibodies, salivary glycoproteins and mucins, which may act as **non-specific blocking agents**.

Once a bacterial cell has successfully attached to a surface in the subgingival area, other host mechanisms come into play.

Desquamation of epithelial cells presents a new cleansing mechanism, which is overcome by certain species by their ability to bind to underlying epithelial cells. Other species are able to invade the epithelial cells and may multiply intracellularly and spread to adjacent cells. Specific **antibody in the subgingival area** could act by **preventing bacterial attachment** or, in some instances, by **making the bacterial cell susceptible to various phagocytic** or killing mechanisms.

A number of subgingival species have evolved mechanisms for evading the effect of specific antibody. Species including *P. gingivalis*, *P. intermedia*, *P. melaninogenica* and *Capnocytophaga* species possess IgG and IgA proteases that can destroy antibody' other species are capable of evading antibody by changing their surface antigens or possibly by mimicking the host's antigens species. A number of bacterial mechanisms exist that might including the production of leukotoxin by *A. actinomycescomitans* and capsules by *P. gingivalis* and other species that inhibit phagocytosis. In addition, a number of species have developed strategies to interfere with the killing mechanisms of the polymorph nuclear leukocytes.

Virulence Factors

Two general mechanisms of pathogenesis have been hypothesized:

- The first involves invasion by subgingival species.
- The second suggests a "long-range" attack where cells of the pathogenic species remain in the pocket but fragments of cells as well as other "**virulence factors**" enter the underlying periodontal tissues and either directly damages the tissues or cause "immune pathology" (indirectly).

Virulence factors can be divided into three categories:

- 1- Substances that damage tissue cells (e.g. H₂S),
- 2- Substances that affect the intercellular matrix (e.g. collagenase). and
- 3- Substances that cause cells to release biologically active substances (e.g. lipopolysaccharide)

Virulence factors of *Aggregatibacter actinomycetemcomitans*

- ❖ **Leukotoxin**; kills PMNs and monocytes.
- ❖ Cytotoxic distending toxin.
- ❖ Immunosuppression factors that inhibit antibody production and activate T-suppressor cells.
- ❖ Inhibition of PMNs functions.
- ❖ Resistant to complement-mediated killing.
- ❖ Lipopolysaccharides.
- ❖ Surface antigens.
- ❖ Antimicrobial resistance.

Virulence factors of *P. gingivalis*

- ❖ Gingipain is a protease secreted by *Porphyromonas gingivalis*. they work to degrade cytokines , thereby down regulating the host response in the form of reduced inflammation.
- ❖ Capsular polysaccharide:
- ❖ Fimbriae, hemagglutinins.
- ❖ Proteinases, hemolysins.
- ❖ Collagenase.
- ❖ secreting a serine phosphatase that inhibits the synthesis of IL-8.
- ❖ Lipopolysaccharides.

Red complex

The red complex is a group of bacteria that are categorized together based on their **association with severe forms of periodontal disease**. The red complex-among a number of other complexes were classified by Sigmund Socransky in 1998.

The three members of the red complex are:

- 1- **Porphyromonas gingivalis.**
- 2- **Tannerella forsythia.**
- 3- **Treponenta denticola.**

NORMAL "CLINICALLY HEALTHY" GINGIVA

Normal gingiva is characterized clinically by its **pink colour** and **firm consistency** and the gingival margin exhibits a **scalloped outline**, The interdental papillae are firm, **do not bleed on gentle probing** and fill the space below the contact areas. The gingiva often exhibits a **stippled appearance** and there is a **knife edge margin** between tooth and soft tissue. Normal gingiva is theoretically free from histological inflammation, but this "ideal" healthy condition has two types: a super healthy or "pristine" state which histologically has little or no inflammatory infiltrate, and the "clinically healthy" gingiva which looks similar clinically but histologically features an inflammatory infiltrate.

In clinically healthy gingiva features an infiltrate of inflammatory cells, **predominantly neutrophils** associated with the junctional epithelium and lymphocytes in the subjacent connective tissue. The infiltrate at this stage may occupy as much as **5%** of the connective tissue volume and is composed of monocytes, macrophages, lymphocytes and neutrophils. These cells are found in the junctional epithelium as well as in the connective tissue of clinically healthy gingiva.



HISTOPATHOLOGICAL FEATURES OF GINGIVAL INFLAMMATION

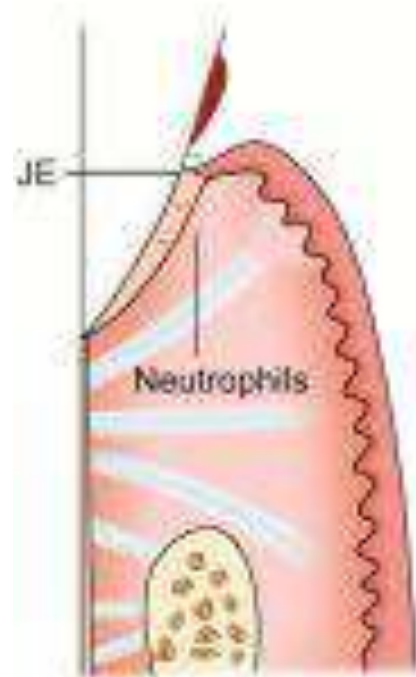
An experimental gingivitis study in dogs was done by Page and Schroeder had compared the cellular and structural composition of the affected area before and during the development of gingivitis over a period of 28 days. At Day 0 of this experiment the normal gingival unit has virtually no inflammatory cells and is comprised of approximately 40-45% epithelium and 55-60 of connective tissue. The connective tissue zone consists of collagen (60%), fibroblasts (13%), vessels (7%) and other tissue constituents, such as intercellular matrix and nerves 20%). Following plaque accumulation, neutrophils and mononuclear leukocytes readily migrate to this area and the connective tissue begins to form and increase in volume over the 28-day period.

At this 28-day interval the connective tissue is comprised of lymphocytes, plasma cells and macrophages which adhere to the collagen matrix and remain in the tissue, whereas neutrophils continue to migrate into the gingival sulcus. With the extensive influx of leukocytes, a marked reduction in the amount of collagen and fibroblasts occurs and the volume of residual tissue (intercellular matrix, degraded collagen, exudates material, degenerated or dead cells) and small blood vessels increases. Page and Schroeder classified the progression of gingival and periodontal inflammation on the basis of **clinical and histopathological evidence into four phases: initial, early, established and advanced** stages or lesions.

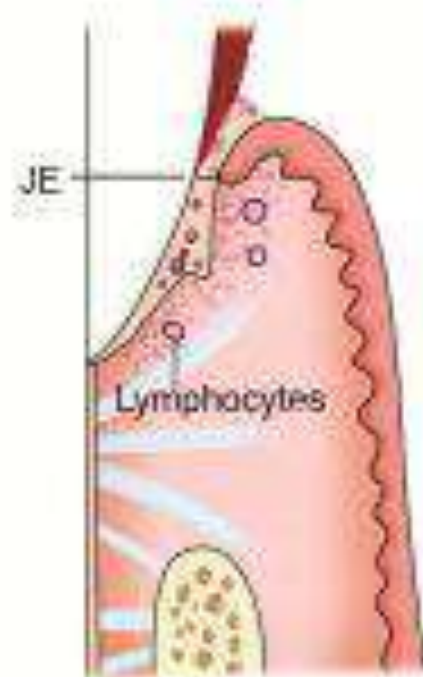
The initial lesion (clinically healthy gingiva)

Inflammation quickly develops as plaque is deposited on the tooth. **Within 24 hours** marked changes are evident in the microvascular plexus beneath the junctional epithelium as more blood is brought to the area. **Dilation of the arterioles, capillaries and venules** of the dentogingival plexus is evident histopathologically. Hydrostatic pressure within the microcirculation increases and intercellular gaps form between adjacent capillary endothelial cells. As the lesion enlarges, and **gingival crevicular fluid flow increases**, noxious substances from microbes will be diluted both in the tissue and the crevice.

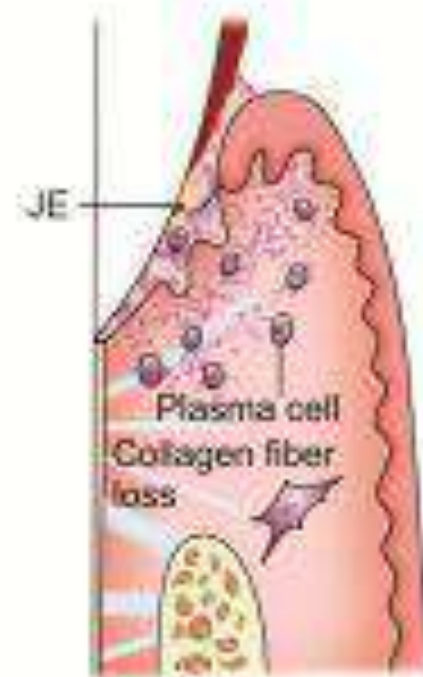
Bacteria and their products may thus be flushed from the sulcus. Plasma proteins escaping from the microcirculation include defensive proteins such as antibodies, complement and protease inhibitors and other macromolecules with numerous functions, probably **within 2-4 days** of plaque build-up the cellular response is well established and is helped by chemotactic substances originating from the plaque microbiota as well as from host cells and secretions. **PMNs** move through the connective tissue and the majority seem to **accumulate in the junctional epithelium and gingival sulcus region.**



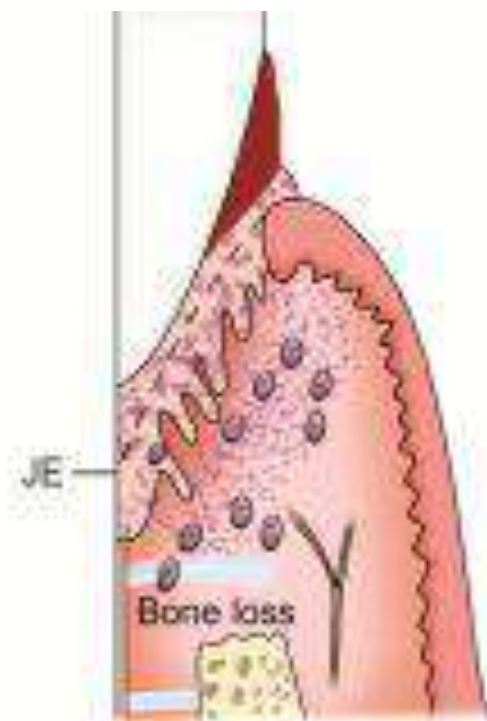
Initial lesion



Early lesion



Established lesion



Advanced lesion
(Bone loss)

The early lesion (early gingivitis)

The early gingival lesion occurs after approximately **one week** of plaque accumulation. **The gingiva is erythematous** in appearance as a result of proliferation of capillaries and continued vasodilatation. Increasing vascular permeability leads to **increased GCF flow**, and transmigrating neutrophils increase significantly in number. The predominant infiltrating cell types are **neutrophils and lymphocytes** (primarily thymic lymphocytes [T-cells]) and the neutrophils migrate through the tissues to the sulcus, and phagocytose bacteria.

Fibroblasts degenerate, primarily via apoptosis (programmed cell death), which increases the space available for infiltrating leukocytes. Collagen destruction occurs, resulting in **collagen depletion** in the areas apical and lateral to the junctional and sulcular epithelium.

The **basal cells** of these epithelial structures **begin to proliferate** to maintain an intact barrier against the bacteria and their products, and as a result the epithelium can be seen proliferating into the collagen depleted areas of the connective tissues. As a result of edema of the gingival tissues, the gingiva may appear slightly swollen, and accordingly, the **gingival sulcus becomes slightly deeper**. thereby **rendering effective plaque control more difficult**. **The early gingival lesion may persist indefinitely or may progress further.**

The established lesion (established gingivitis)

Generally there is a further enhancement of the inflammatory state as exposure to plaque continues. There is **increased fluid exudation and leukocyte migration** into the tissues and the gingival crevice. Clinically this lesion will exhibit **more edematous swelling than the "early gingivitis"**.

Plasma cells are seen situated primarily in the coronal connective tissues and the **rete pegs extend deeper** into the connective tissue in an attempt to maintain epithelial integrity and a barrier to microbial entry.

The pocket epithelium that now has formed has a heavy leukocyte infiltrate, predominantly of PMNs which eventually migrate across the epithelium into the gingival pocket. In comparison to the original junctional epithelium, the pocket epithelium is more permeable to the passage of substances into and out of the underlying connective tissues and may in places be temporarily ulcerated. The pocket epithelium is less able to resist the passage of the periodontal probe, so **bleeding on probing is a common feature** of chronic gingivitis. It is important to remember that these inflammatory changes are still completely **reversible** if effective plaque control is reinstated.

The advanced lesion (periodontitis)

The advanced lesion marks the **transition from gingivitis to periodontitis**.

This transition is determined by many factors, include the bacterial challenge (both the composition and the quantity of the biofilm), the host inflammatory response, and susceptibility, factors, including environmental and genetic risk factors.

So the final stage in this process is known as the advanced lesion. The lesion is no longer localized to the gingival, and **the inflammatory cell infiltrate extends laterally and apically into the connective tissue of the true attachment apparatus**. The advanced lesion has all the characteristics of the established lesion but differs importantly in that **alveolar bone loss** occurs, fiber damage is extensive, **the junctional epithelium migrates apically from the cemento-enamel junction**, and there are widespread manifestations of inflammatory and immunopathological tissue damage. It is generally accepted that **plasma cells are the dominant cell type** in the advanced lesion.

Thank you

Periodontal management of medically compromised patients

Many patients seeking dental care have significant medical conditions that may alter both the course of their oral diseases and the therapy provided. The older age of the periodontal patients increases the likelihood of underlying diseases. Therefore therapeutic responsibility of the clinician includes identification of the patients medical problem to formulate proper treatment plans. Thorough medical histories are important and sometimes consultation with or referral of the patients to an appropriate physician may be indicated. This ensures correct patients management and provided medico-legal coverage to the clinician.

From the most common medical problems are the following:

1-cardio-vascular diseases:-

These diseases are the most prevalent category of systemic diseases and more common with increasing age. They include hypertension, angina pectoris, myocardial infarction, previous cerebrovascular accident, congestive heart failure, presence of cardiac pacemaker and infective endocarditis. In most cases the patient physician should be consulted, especially if stressful or prolonged treatment is anticipated. Short appointment and a calm , relaxing environment help to minimize stress.

a)Hypertension:

It is the most common cardiovascular disease and it is defined as a systolic blood pressure of 140 mmHg or greater, or a diastolic blood pressure of 90 mmHg or greater , and it is diagnosed on a single elevated blood pressure recording but it is based on the average value of three or more blood pressure readings taking at three or more appointment. If hypertension persist and increase in severity, it may lead to coronary heart disease, angina, congestive heart failure , cerebrovascular accident or kidney failure. Management of those patients will be based as follows:

- 1-No periodontal treatment should be given to patients who is hypertensive and not under the medical management.
- 2-The dentist should inform the physician about the degree of stress, blood loss and length of the periodontal procedure so that avoid excessive bleeding.

3-Local anesthesia without epinephrine may be used for short procedure (less than 30 minute)or use local anesthesia with an epinephrine concentration not more than 1:100:000 to control pain and minimize stress (dental treatment for hypertensive patients is generally safe as long as stress is minimized.

4-Avoid sudden positional changes or syncope.

b) Angina pectoris:

Angina occurs when myocardial oxygen demand exceed supply, resulting in temporary myocardial ischemia. Patients with a history of unstable angina pectoris (angina that occurs irregularly or on multiple occasions without predisposing factors) should be treated for emergencies only and in consultation with the patient physician. Patient with a history of stable (angina that is associated with stress and easily controlled with medication and rest) can be treated with the following precautions:

1-Premedication if needed as valium.

2-Morning and short appointment.

3-Nitroglycerine medication sublingually 5 minute before the procedure.

4-If during the periodontal procedure , the patient become fatigue or uncomfortable , the procedure should be discontinued.

c)Cardiac pacemaker:

Some cardiac arrhythmias are treated with implanted pacemakers which usually implanted in the chest wall and enter the heart transvenously. These electrical devices are used to regulate heart beats and no electro-physiologic problems may occur with such implanted device. Managements of such patients will be as follows:

1- Consult with the physician to get information about the underlying cardiac reason for pacing and to explain the periodontal treatment plan to him.

2-The patients should be positioned so that minimal pressure will be exerted on the implanted site.

3-Limited use of electrical equipment that generated electromagnetic field such as ultrasonic devices so that to avoid interferences with the artificial pacemaker. Try to keep these device at least 30 cm from the patient. However, most pacemakers are adequately to prevent these changes.

d) Infective endocarditis

It is a diseases in which microorganisms colonize the damaged endocardium or heart valve . it is a serious diseases with poor prognosis. The term infective

endocarditis is preferred to the previous term bacterial endocarditis because the disease can also be caused by fungi and viruses. The organisms most commonly encountered in IE are a hemolytic streptococci (streptococcus viridans) . other m.o. found in the periodontal pocket and associated with this disease are Eikenella corrodens, A.a., capnocytophaga and lactobacillus species. The practice of periodontics is intimately concerned with the prevention of IE. Any dental procedures that involve bleeding may induce a transient bacteremia, so prophylactic antibiotic should be recommended before the procedures which is associated with significant bleeding as periodontal surgery, scaling and root planning. However , bacteremia may occur even in the absence dental procedures, especially in individuals with poor hygiene and significant periodontal information. The preventive measures to reduce the risk of IE should consists of following:

- 1-Define the susceptible patients: those patients at high risk to develop IE following dental treatment include those with rheumatic heart diseases , congenital heart diseases, cardiac surgery , prosthetic heart valves.
- 2-Provide oral hygiene instruction: in patients with significant gingival inflammation, oral hygiene should be initially limited to gentle procedures (oral rinses as chlorhexidin mouth rinse and gentle tooth brushing with soft brush). As gingival health improves, more aggressive oral hygiene may be initiated.
- 3-During periodontal treatment, recommended prophylactic antibiotic regimens should be practiced with all susceptible patients. The regiment used is the following:

Regimen	Antibiotic	Dosage
Standard oral regimen	amoxicillin	2 g 1 hour before procedure

Alternate regimen for patients allergic to amoxicillin or penicillin	Clindamycin or	600mg 1 hour before procedure
	azithromycin	500mg 1 hour before procedure
Patient unable to take oral medication	ampicillin	2g intramuscularly or intravenously within 30 minute before procedure
Patients unable to take oral medication and allergic to penicillin	clindamycin	600mg intravenously within 30 minute before procedure (must be diluted and injected slowly)

Patients with aggressive periodontitis often have high level of Aggrigatibacteractinomycetemcomitans in the sub-gingival plaque. This organism has been associated with IE and is often resistant to penicillin , therefore in patients with aggressive periodontitis who are also at risk for IE, it has been suggested using tetracycline 250mg four times daily for 14 days to eliminate or reduce A.a. , followed by the conventional prophylaxis protocol at the time of dental treatment.

4) Periodontal treatment should be designed according to the degree of severity and involvement of periodontal tissues:- periodontal therapy is prolonged procedure , it is mostly not a one day antibiotic regimen, multiple visits and easily elicit gingival bleeding , so periodontal treatment plans must be developed for patients susceptible to IE and as follows:

a- In order to reduce the wide range systemic effect of periodontal diseases in these patients, teeth with severe periodontitis and poor prognosis have to be extracted rather than retained and treated.

b- All periodontal treatment procedures (including probing) require antibiotic prophylaxis. Pretreatment chlorhexidine mouth rinse are recommended before all procedure because it reduce the presence of bacteria on mucosal surfaces.

c- Reduce the number of visits required so that to minimize the risk of developing resistant bacteria.

d- It is preferably that the appointments allowed between 10-14 days, if it is not possible then select an alternative antibiotic regimen.

e- The need for antibiotic prophylaxis before suture removal is controversial when possible use the resorbable suture in such patients.

f- Regular recall appointment are important with reinforcement on good oral hygiene to maintain periodontal health.

2- Renal disease

Patients with chronic renal failure have a progressive diseases that may require kidney transplantation or dialysis. The patients who are receiving hemodialysis require special precautions. Those patients have a high incidence of viral hepatitis, anemia, and prolonged hemorrhage. The risk of hemorrhage is related anticoagulants during dialysis. Also they have either an internal arteriovenous fistula or an external arteriovenous shunt. This shunt is often located in the arm and must be protected from trauma.

The management of those patients will be as follows:

1-Consult with patient physician.

2-Screen for hepatitis B surface antigen and antibodies prior to any treatment.

3-Avoid drugs that metabolized by the kidney ex. Tetracycline , streptomycin, aminoglycoside, aspirin.....

4-Provide antibiotic prophylaxis to prevent infective endocarditis .

5-Screen for bleeding disorder(bleeding time and platelets time)
(normally bleeding time= 1-6 seconds, platelets =140,000-400000/mm)

6- Monitor blood pressure because those patients usually hypertensive.

Patients with renal transplantation take immunosuppressive drugs that greatly reduce resistance to infection, So management of those patients will be as follows:

1-Prophylactic antibiotic to prevent infection (prescribed by the physician)

2- May need supplemental corticosteroid .

3-Teeth with furcation involvement , periodontal abscesses should be extracted if it is not savable before transplantation to reduce possibility of infection.

4-Surgical excision of the gingiva may be needed because of gingival overgrowth secondary to cyclosporine therapy.

3-Endocrine disorders

a)Diabetes Mellitus

The diabetic patients require special precautions before periodontal therapy. The two major types of diabetic type 1 (formerly known as insulin-dependent diabetics) and type 2 (formerly called non-insulin dependent diabetic). Diabetic patients are managing their blood glucose levels (glycaemia) through diet, oral agent and insulin therapy. The classic signs of diabetic include polydipsia (excessive thirst). Polyuria (excessive urination), and polyphagia (excessive hunger with unexplained weight loss). If the patients has any of these signs and symptoms , physician consultation is indicated for further investigation because periodontal therapy has limited success in the presences of undiagnosed or poorly controlled diabetic. If the patients is suspected of having undiagnosed diabetic , following procedures should be performed:

1-Consult patients physician

2-Analyze laboratory tests

a) Fasting blood glucose ≥ 126 mg/dl. Fasting is defined as no caloric intake for at least 8 hours (normal fasting glucose 70-100mg/dl).

b) Symptoms of diabetic plus non fasting plasma glucose ≥ 200 mg/dl. Non fasting glucose may be drawn at any time of the day without regard to time since the least meal.

c) 2 hours postprandial glucose tolerance test. (the glucose level is measured immediately before and 2 hours after a person drinks a liquid containing 65 g of glucose dissolved in water). Normal 2 hours postprandial glucose is ≤ 140 mg/dl.

3- Provide emergency periodontal therapy only for such patient like acute periodontal abscess until diagnosed is established.

If a patient is known to have diabetic, it is important to determine the level of glycemic control before initiating periodontal treatment. The primary test used to assess glycemic control in a known diabetic individual is the glycosylated or glycated hemoglobin test (HbA1c) is a fraction of hemoglobin that reflected blood glucose concentration over the preceding 6 to 8 weeks and may provide an indication of the potential response to periodontal therapy. Well-controlled diabetic patient (HbA1c $\leq 8.5\%$) usually respond to therapy in a manner similar to non diabetic individuals. Poorly controlled patients (HbA1c $\geq 8.5\%$) often have a poor response to treatment, with more postoperative complications.

b) Thyroid disorders:

Hyperthyroidism or thyrotoxicosis may increase risk for hypertension, angina, congestive heart failure. So

1-Avoid any periodontal treatment for patients with thyrotoxicosis until good medical control.

2-Avoid epinephrine in completely treated patient.

3-Avoid stress and control periodontal infection to prevent the occurrence of thyrotoxic crisis in untreated patient.

4- Once under good medical management, patient may receive dental treatment.

In Hypothyroidism

1-Avoid stress and infection to prevent the occurrence of hypothyroid coma.

2-Avoid narcotic and tranquilizer in untreated hypothyroid patients because of inability to tolerance drugs.

3-In patients under good medical management, dental treatment may be performed.

4) Pregnancy

The aim of periodontal therapy for pregnant woman is to reduce the exaggerated inflammatory response of the periodontal tissues to local factor which related to hormonal changes associated with pregnancy.

-The second trimester is the safest time for treatment (scaling. Polishing, root planning) while surgical procedures should be postponed after delivery.

-In the third trimester, treatment should is not advisable because of supine hypotension syndrome of pregnancy , loss of consciousness may occur due to pressure of the uterus on inferior vena cava.

-No medication is given that the placenta and affect the fetus.

-No radiograph unless necessary with persuasions.

5) Hemorrhagic disorders:

Patients with a history of bleeding problems caused by disease or drug should be managed to minimize risks of hemorrhage. Identification of these patients via the health history. Clinical examination and clinical laboratory tests is important. Health questioning should cover

1-History of bleeding after previous surgery or trauma.

2-Past and present drug history.

3-History of bleeding problems among relatives.

4-Illnesses associated with potential bleeding problems.

Clinical examination should detect the existence of jaundice, ecchymosis, petechial, spontaneous gingival bleeding.

Laboratory tests include bleeding time, prothrombin time, complete blood count, partial thromboplastin time and coagulation time.

Bleeding disorders may include the following:

- 1-Hemophilia **A** (result in a deficiency of coagulation factor VIII).
- 2-Hemophilia **B** (result in a deficiency of factor IX).
- 3- Von Wille brand disease (result from deficiency of Von willebrand factor which mediates adhesion of platelet to the injured vessels wall).
- 4- Liver disease: Most coagulation factor are synthesized by the liver or it is the site for production of the clotting factors. Long term alcohol abusers or chronic hepatitis patients often demonstrated inadequate coagulation.
- 5-Patients taking anti-coagulant drugs: patients with prosthetic heart valves, or histories of myocardial infraction, stroke or thromboembolism are frequently placed on anticoagulation therapy using dicumarol and warfarin. These drugs are vitamin K antagonists.

Another drug is aspirin which interferes with normal platelete aggregation and can result in prolonged bleeding. For patients taking more than 325mg of aspirin per day, the drug should be discontinued at least 7 to 10 days before periodontal therapy in consultation with the physician.

6-Thrombocytopenia purpuras

Thrombocytopenia is defined as a platelet count ≤ 100.000 mm. the purpuras could result from radiation, chemotherapy , leukemia, or infection and it is characterized by spontaneous petechiae (small red patches)or ecchymosis.

In general speaking for patients with bleeding disorders, never do any type of periodontal treatment unless consultation with the physician and it is preferable to do periodontal surgery if needed in the hospital.

8) Infectious diseases:

Because medical histories are often inaccurate or incomplete, all periodontal patients should be treated as if they have infectious diseases. Protection of patients, clinicians and office staff requires use of universal (standard) precautions for each patient. An example of these diseases are hepatitis, AIDS, and Tuberculosis.

AIDS:

AIDS is characterized by impairment of the immune system. The human immunodeficiency virus (HIV) was isolated in 1984 as the causative agent or virus of AIDS. Most of the patients develop long lasting acute illness with flu-like symptoms that last for 10-14 days with enlarged lymph nodes, night sweating, weight loss, fever, malaise, and chronic diarrhea. Oral manifestations characterized by oral hairy leukoplakia and oral candidiasis, necrotizing ulcerative gingivitis or periodontitis (NUG OR NUP). Periodontal management of AIDS patients involves.

- Using full barrier technique.
- Care in use of all sharp instruments.
- Proper sterilization.
- Do not use ultrasonic instrumentation.

Tuberculosis

The patients with tuberculosis should receive emergency care only. Physician should be consulted for the result of sputum cultures for Mycobacterium tuberculosis. When the results are negative, the patients may be treated normally. When the results are positive – we have to know that adequate treatment of tuberculosis requires a minimum of 18 months with a post-treatment follow-up. So periodontal treatment should include emergency only. In general, in case of infectious diseases it is preferable to wear double gloves and double masks. The sterilization should be done in an autoclave at 120-130°C for about one to two hours.

Hepatitis

Six distinct viruses causing viral hepatitis have been identified A,B,C,D,E, AND G viruses. These forms differ in their virology, epidemiology, and prophylaxis.

1-Hepatitis A and E are both self limiting infection with no associated chronic liver disease and these viruses transmitted via fecal-oral route.

2-Hepatitis B infection may result in chronic liver diseases, it transmitted mainly through hematogenous routes and through contaminated instruments or needle in the dental office. Hepatitis B vaccine is recommended for all care health workers.

3-Hepatitis D viruses require the presence of hepatitis B virus to survive and replicate because the virus genetic material is package within the hepatitis B virus surface antigen coating. So prevention of this virus depends strongly on hepatitis B virus vaccination.

4-Hepatitis C is the most serious infection due to high chronic infection rate. Only 15% of patients infected with this virus recover completely and 85% develop chronic infection which increase the risk for cirrhosis and liver failure. No vaccine is available for this virus.

5-Hepatitis G is a newly discovered virus and it is virology is not clearly understood and it known to be transmitted via blood.

> for patients with past history of hepatitis, consult the physician to determine the type of hepatitis, course, length of the disease and mode of the transmission.

> if the disease is in the active stage, do not provide periodontal treatment.

> for recovered type A or E hepatitis patients, perform routine periodontal care.

> for recovered type B and D hepatitis you must screen for HBsAG. If this test is positive , so the patient is infective.

> patient with positive anti-HBs may be treated routinely.

> patients with active hepatitis and need emergency treatment, we should do the following:

1-using full barrier techniques including masks, gloves, and eye glasses.

2-do not use ultrasonic instrument or air syringe so that not to transfer the infection by the saliva.

3-rinsing with chlorhexidin mouth wash is recommended.

4-when the procedure is complete, all instruments should be sterilized carefully.

PERIODONTAL DISEASE AND DIET

اعداد:

د. نور صباح ارحيم



PALEO
LEAP

- **Periodontal health** is influenced by a number of factors such as **oral hygiene**, **genetic** and **epigenetic** factors, **systemic health**, and **nutrition**. Many studies have observed that a balanced diet has an essential role in maintaining periodontal health. Additionally, the influences of nutritional supplements and dietary components have been known to affect healing after periodontal surgery. Studies have attempted to find a correlation between tooth loss, periodontal health, and nutrition. Moreover, bone formation and periodontal regeneration are also affected by numerous vitamins, minerals, and trace elements.

-Clinicians often overlook the effect of nutrition on the **immune system** and its role in periodontal disease progression. It is an important consideration, as we understand that diet plays a modifying role in the progression of periodontal disease..

-A variety of nutrients have a major impact on periodontal health. Nutrients are of two types: **micronutrients and macronutrients**. **Micronutrients** are those components of food that are required in small or trace amounts. The human diet contains a number of antioxidants in the form of micronutrients .

-Antioxidant micronutrients include vitamin **A** (carotenoids and -carotene), vitamin **C** (ascorbic acid) , vitamin **E** (-tocopherol) , glutathione , and melatonin .

. ▶

Macronutrients are nutrients required in large quantities, for example minerals, proteins, carbohydrates, and fats in addition to oxygen and water. High carbohydrate intake has been implicated in periodontal disease and dental caries . The relationship of nutrition and oral health is well known . For example, a sugary diet encourages plaque formation and leads to the onset or worsening of dental decay in reaction to poor oral hygiene

Dietary supplements in the treatment of periodontal disease: ▶

1-The role of vitamin **C** in periodontitis ▶

2-A low dietary intake of **calcium** is associated with severe periodontal attachment loss, and prevalence of periodontal disease decreases with high intake of dairy products ▶

3-**B**-vitamin supplementation results in higher clinical attachment following flap ▶
Surgery. ▶

4-Vitamin **D** deficiency contributes to negative outcomes following periodontal surgery ▶

5- There is a positive correlation between Vitamin **D3** supplementation and ▶
osseointegration of dental implants . ▶

6-A higher intake of vitamins **A, B, C**, and E along with omega-3 fatty acids results in improved healing after ▶
non-surgical periodontal therapy.

7-Vitamin K is a group of vitamins required for the synthesis of proteins that are precursors or ▶
prerequisites of the formation of blood coagulation factors such as prothrombin and factors VII, IX, and ▶
X [71]. In addition, research has indicated that vitamin K also plays a role in the formation of proteins ▶
required for bone metabolism such as osteocalcin and periostin ▶

8-Probiotics are defined by the World Health Organization as "live microorganisms, which, when administered in adequate amounts, confer a health benefit on the host." Also known as "good bacteria," the probiotics most well-known are Lactobacillus and Bifidobacterium, each of which has several strains. Most common sources of probiotics include yogurts, select supplements, lozenges, juices, milks, cheeses, and soy products.

9-another interesting finding is that **Omega-3** (ν -3) polyunsaturated fatty acids (PUFA), including docosahexaenoic acid (DHA) and eicosapentaenoic acid (EPA), were shown to have therapeutic anti-inflammatory and protective actions in inflammatory diseases including periodontitis.

10-The production of **lactic acid** through carbohydrate fermentation by oral lactobacilli generates a low pH, which might inhibit the growth of anaerobic bacteria. One of the important differences between lactic acid foods and other dairy products, such as milk and cheese, is the presence of lactobacilli. The plausible hypothesis that lactic acid foods may have a beneficial effect on periodontal disease might be based on the probiotic effect of lactobacilli in these foods. The regular consumption of lactic acid foods may constrain periodontal disease by controlling the overgrowth of periodontal pathogens in the oral cavity.

1-calcium: Calcium is required for the normal functioning of muscles and body systems. Additionally, calcium is essential for the maintenance and formation of calcified tissues such as bone and teeth.

2-Magnesium: Magnesium is required for cell metabolism and maintenance and formation of bone. The deficiency of magnesium interfered with the parathyroid hormone and directly affects the bone resulting in osteoporosis.

3-Iron: Iron is mainly required for synthesis of proteins, including hemoglobin and enzymes. Foods such as red meat, spinach, fish (tuna and salmon), and beans are rich sources of iron. Iron deficiency leads to anemia and related symptoms. Oral manifestations of anemia include recurrent ulceration, pale mucosa, and burning of the mouth.

4-Zinc: Protein-rich foods are the primary source of dietary zinc . Zinc is second to iron as the most abundantly found trace mineral in the human body . Zinc acts as a cofactor in many enzyme-controlled processes. Particularly, it modulates the processes of auto-debridement and keratinocyte migration during wound repair

5-Fluoride: The anti-caries effects of fluoride have been established for a long time . Fluorine prevents

caries by strengthening enamel and cementum due to the formation of fluoroapatite and exerting an antibacterial effect via inhibition of bacterial growth and adhesion . Hence, topical fluoride, in the form of dentifrices, gels, foams, and varnishes has been used as a preventative measure against dental caries . Considering its beneficial roles, fluoride has been incorporated into various restorative materials such as glass ionomers . These materials act as reservoirs that are capable of releasing fluoride into the oral cavity and recharging while fluoride is available from toothpaste, mouthwashes, or fluoride-rich foods. Systemic administration of fluoride may be via water, milk, and capsules. ▶

6-Melatonin is a potent antioxidant secreted by various organs of the human body . Additionally, plants and cereals are sources of melatonin. Although melatonin is not classified as a major nutrient, it has been suggested that, in supplement form, the antioxidative properties of melatonin are more potent than those of Vitamin E: ▶

7-Lycopene is a red pigment present in vegetables such as tomatoes, carrots, and watermelons. ▶

Lycopene may prevent cancer and cardiac diseases due to its antioxidant effects [103]. Similarly, in ▶
some studies lycopene has been investigated as an adjunct to non-surgical periodontal therapy. A ▶
study by Chandra et al. suggested that lycopene supplementation may enhance the improvement of ▶
periodontal health [10 ▶

The 12 Best Foods to Eat if You Have Gum Disease

1. Nuts and seeds high in omega 3s

These include macadamia nuts, pistachios, sesame seeds. Omega-3s are essential to a well-rounded, nutrient-dense diet. Omega-3s are great for you because they are anti-inflammatory but also have an inverse relationship with gum disease. These nuts are great sources of nutrients as well.

2. Salmon or other fatty fish

Oily fish such as salmon, herring, and mackerel are great sources of nutrients and are high in omega-3s. Due to their oil content, they have anti-inflammatory effects on the body and are great for the immune system.

3. Grass-fed beef

Because we don't eat organ meats like our ancestors did, we need to be conscientious about our collagen intake. Grass-fed beef contains collagen, which is great for fighting gum disease.

It's also high in omega-3s, which is one of the reasons so many health gurus specify the feed of the cow. Grain fed beef is packed with omega-6s, antibiotics, and other inflammatory, disease causing components. It's just not worth it to eat grain-fed beef anymore.

- ▶ **4. Chicken** :Chicken contains CoQ10 and collagen, which are both excellent in the fight against gum disease. Don't skip the skin either. Many people forgo eating the chicken skin because it's high in fat, but chicken skin is packed with collagen.
- ▶ **5. Bone broth**: We don't eat the whole animal like we once did, which means we're missing out on important nutrients like collagen. But you can get more of this nutrient in your diet with bone broth. Bone broth is gaining popularity as a health superfood because it's a delicious way for us to get more of the nutrients we miss out on in the modern world.
- ▶ **6. Shiitake mushrooms**: Shiitake mushrooms contain lentinan, which is a polysaccharide only found in this particular type of mushroom. Lentinan attacks the harmful gum disease causing bacteria while leaving other bacteria alone. Shiitake mushrooms are also powerfully anti-inflammatory and have been seen in studies to reduce numerous inflammatory markers.
- ▶ **7. Broccoli** :Broccoli is packed with Vitamin C which acts as an antioxidant in the body. Vitamin C has been inversely associated with gum disease. Broccoli also provides tons of other nutrients. Broccoli contains Vitamin B1, magnesium, iron, calcium, niacin, and selenium. Any nutrient dense diet should contain broccoli.
- ▶ **8. Red and green bell peppers**: High in Vitamin C, red and green bell peppers are a great addition to a gum disease fighting diet. One cup of red bell peppers contains over 300% of your daily recommended allowance of Vitamin C. While green bell peppers contain over 200% of the Vitamin C you need in a day. In fact, both red and green bell peppers have more Vitamin C than an orange. Add more bell peppers to your diet for an easy and delicious boost in Vitamin C.
- ▶ **9. Sweet potatoes** :Sweet potatoes are an excellent source of many nutrients but they made this list specifically for their beta carotene content. Beta carotene is an essential nutrient in the fight against gum disease. It reduces inflammation and provides your body with the elements it needs to create Vitamin A. Sweet potatoes also contain high levels of Vitamin C, Vitamin B6, and Manganese.

- ▶ **10. Green tea** Green tea is high in catechins. Remember catechins? Those were one of the key gum disease fighting nutrients listed above. Not only is green tea high in catechins, but it's also been found to have an inverse relationship with periodontal disease. One study that analyzed 940 men found there was an inverse association between green tea consumption and periodontal disease. Meaning those that drink more green tea have a lower chance of developing gum disease.
- ▶ **11. Cacao** Cacao is also high in disease-fighting catechins, which help stop gum disease. Cacao is great for your teeth overall because it fights cavities, plaque, and tooth decay . Compounds in cacao are more effective at fighting tooth decay than fluoride. Cacao also contains a compound called CBH, which hardens your enamel. To eat cacao that's actually good for you and not loaded with sugar, go for a raw chocolate that is 70 percent or more. Look for a low sugar content too. Cacao and cocoa differ in processing. Cacao hasn't been exposed to high heat, is still raw, and still contains it's 300+ compounds, many of which are beneficial to your health.
- ▶ **12. Probiotic containing foods** These include kefir, sauerkraut, and kimchi. Probiotics support the good bacteria in your mouth and reduce gingivitis and plaque buildup. Fermented foods are also thought to suppress the growth of oral pathogens. One study found those who drank fermented dairy benefited from protective effects against gum disease – though it wasn't clear whether it was due to the increase calcium or the probiotics (or both!).

Helping our patients understand ►

In treating the periodontal patient successfully, how many of us stop to consider the role of the patient's diet in the healing process?

Having a patient keep a weekly food diary was a task we completed in hygiene or dental school, but few clinicians incorporate serious nutritional counseling into the treatment protocol.

THANK YOU

Periodontal indices



Plaque index (Silness and Loe 1964)

- ▶ **This index measures the thickness of plaque on the gingival one third of the teeth.**
- ▶ used to evaluate the level and rate of plaque formation on tooth surfaces, and to test the efficacy of oral care products for removal and prevention of plaque deposits from these surfaces.
- ▶ Used on all teeth (28, wisdom teeth are excluded) or selected teeth (6 teeth) .
- ▶ Used on all surfaces (4) (M, B, D, L).
- ▶ • **Score 0** No plaque
- ▶ • **Score 1** A film of plaque adhering to the free gingival margin and adjacent area of the tooth, which can not be seen with the naked eye. But only by using disclosing solution or by using probe.
- ▶ • **Score 2** Moderate accumulation of deposits within the gingival pocket, on the gingival margin and/ or adjacent tooth surface, which can be seen with the naked eye.
- ▶ • **Score 3** Abundance of soft matter within the gingival pocket and/or on the tooth and gingival margin.

Gingival Index (GI) (Loe and Silness, 1967)

- ▶ measures the degree of gingival inflammation. Tissues surrounding each tooth divided into 4 gingival scoring units: distal facial papilla, facial margin, mesial facial papilla, lingual gingival margin.
- ▶ **Score of gingival index**
- ▶ **Score 0** Normal gingiva
- ▶ **Score 1** Mild inflammation — slight change in color, slight edema. No bleeding on probing
- ▶ **Score 2** Moderate inflammation — redness, edema and glazing. Bleeding on probing
- ▶ **Score 3** Severe inflammation — marked redness and edema. Ulceration. Tendency to spontaneous bleeding.

The GI may be used for the assessment of prevalence and severity of gingivitis in populations, groups and individuals.

Plaque index (O'leary)

- a commonly used oral hygiene index for assessing oral health skills. This index provides sufficient information for patient education.
- Suitable disclosing solution is painted on all exposed tooth surfaces.
- The operator (using an explorer or a tip of a probe) examined each stained surface for soft accumulations at the dentogingival junction. When found, they are recorded by making a dash/red colour in the appropriate spaces on the record form.

Plaque index (Quigley Hein)

Score

- 0 No plaque at the cervical margin
- 1 Separate flecks of plaque at the cervical margin of the tooth
- 2 A thin continuous band of plaque (≤ 1 mm) at the cervical margin of the tooth
- 3 A band of plaque wider than 1 mm but covering less than one-third of the crown of the tooth
- 4 Plaque covering at least one-third but less than two-thirds of the crown of the tooth
- 5 Plaque covering two-thirds or more of the crown of the tooth

Example

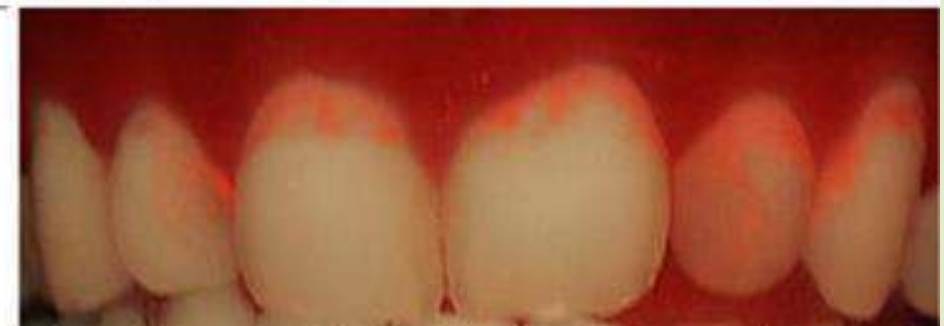
Blue-mQH

1 2 3 3 4 2



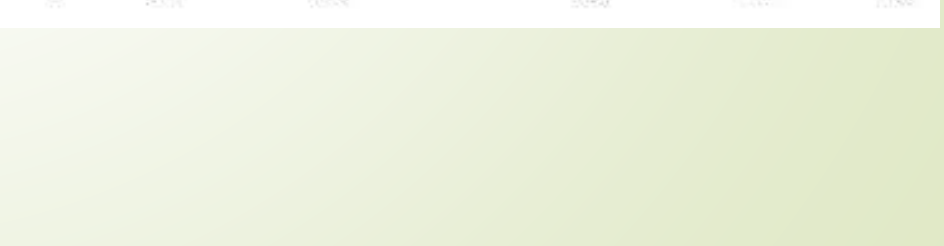
Combi-mQH

2 5 4 4 5 3



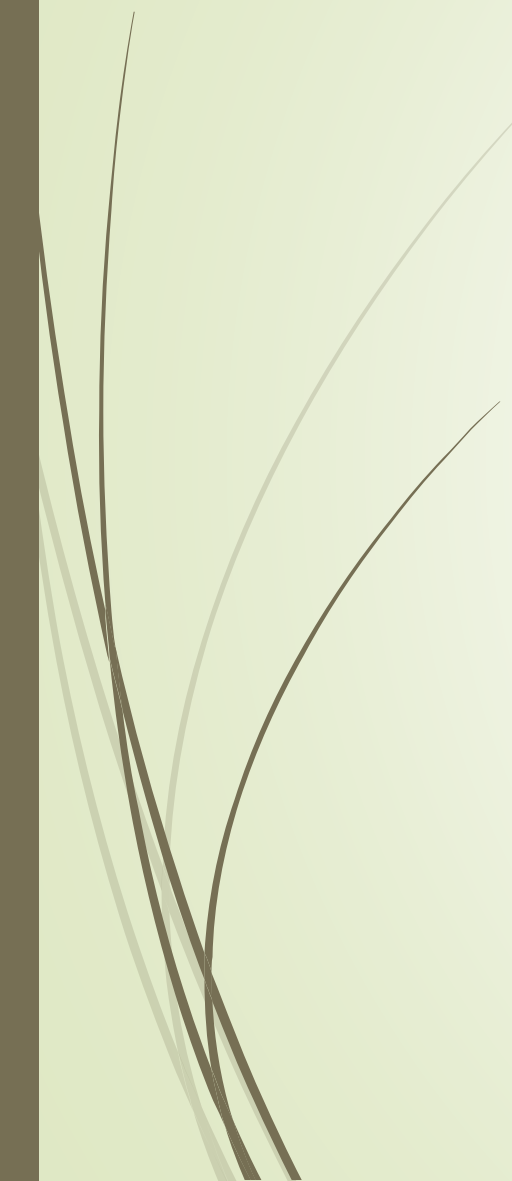
QLF-mQH

1 3 3 3 4 3





Bleeding on probing (BOP)

- ▶ A periodontal probe is inserted to the —bottom of the gingival/periodontal pocket by applying light force and is moved gently along the tooth (root) surface. If bleeding is provoked upon retrieval of the probe, the site examined is considered —BoP — positive and, hence, is inflamed
- 

Calculus Index (CI)

- Calculus is mineralized material on the tooth surface.
- The calculus index refers to the amount of calculus on a tooth.
- **CI 0** — No observable calculus.
- **CI 1** — Supragingival calculus covering not more than 1/3 of the exposed tooth surface.
- **CI 2** — Supragingival calculus covering more than 1/3 but not more than 2/3 of the exposed tooth surface or presence of flecks of subgingival calculus.
- **CI 3** — Supragingival calculus covering more than two-thirds of the exposed tooth surface or a continuous heavy band of subgingival calculus around the cervical portion of the tooth.

Probing pocket depth

- The probing depth, that is the distance from the gingival margin to the bottom of the gingival sulcus/pocket, is measured to the nearest millimetre by means of a graduated periodontal probe.
- The probe should be inserted parallel to the vertical axis of the tooth and “walked” circumferentially around each surface of each tooth to detect the areas of deepest penetration. This turn means that single-rooted teeth have to be examined at four sites at least (e. g. mesial, buccal, distal, and oral) and multirouted teeth at six sites at least (e. g. mesiobuccal, buccal, distobuccal, distooral, oral, and mesio-oral)

Clinical attachment loss (CAL)

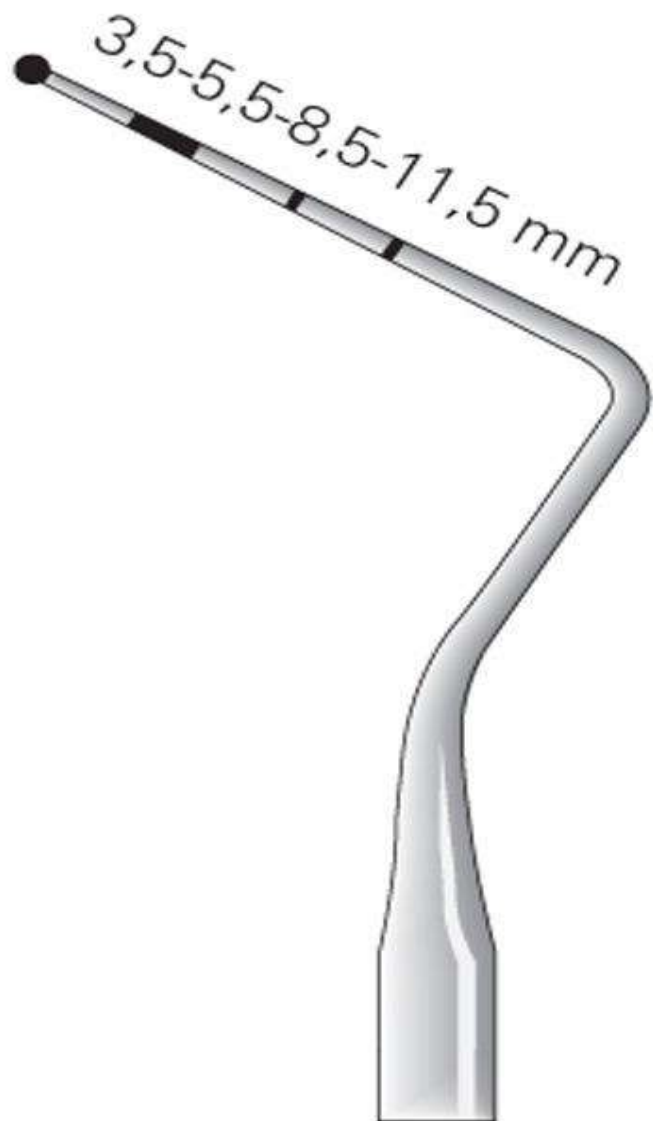
- is a more accurate indicator of the periodontal support around the tooth than probing depth alone.
- CAL is measured from a fixed point on the tooth that doesn't change, the CEJ.
- To calculate CAL, two measurements are needed:
- **1-In recession** : probing depth + gingival margin to the CEJ (add)
- **2- In tissue overgrowth** : probing depth – gingival margin to the CEJ(subtract)

BASIC PERIODONTAL EXAMINATION (BPE)

- The BPE is a simple and rapid screening tool that is used to indicate the level of examination needed and to provide basic guidance on treatment need. Please note; the **BPE does not provide a diagnosis.**
- 1. The dentition is divided into 6 sextants:
 - upper right (17 to 14), upper anterior (13 to 23), upper left (24 to 27)
 - lower right (47 to 44), lower anterior (43 to 33), lower left (34 to 37)
- 2. All teeth in each sextant are examined (with the exception of 3rd molars).
- 3. For a sextant to qualify for recording, it must contain at least 2 teeth. (If only 1 tooth is present in a sextant, the score for that tooth is included in the recording for the adjoining sextant).

BASIC PERIODONTAL EXAMINATION (BPE)

- 4. A WHO BPE probe is used (World Health Organisation probe). This has a “ball end” 0.5 mm in diameter, and a black band from 3.5 to 5.5 mm. Light probing force should be used (20-25 grams).
- 5. The probe should be “walked around” the sulcus/pockets in each sextant, and the highest score recorded. As soon as a code 4 is identified in a sextant, the clinician may then move directly on to the next sextant, though it is better to continue to examine all sites in the sextant. This will help to gain a fuller understanding of the periodontal condition, and will make sure that furcation involvements are not missed. If a code 4 is not detected, then all sites should be examined to ensure that the highest score in the sextant is recorded before moving on to the next sextant.



WHO BPE probe

BASIC PERIODONTAL EXAMINATION (BPE)

Scoring codes

0	No pockets >3.5 mm, no calculus/overhangs, no bleeding after probing (<i>black band completely visible</i>)
1	No pockets >3.5 mm, no calculus/overhangs, but bleeding after probing (<i>black band completely visible</i>)
2	No pockets >3.5 mm, but supra- or subgingival calculus/overhangs (<i>black band completely visible</i>)
3	Probing depth 3.5-5.5 mm (<i>black band partially visible, indicating pocket of 4-5 mm</i>)
4	Probing depth >5.5 mm (<i>black band entirely within the pocket, indicating pocket of 6 mm or more</i>)
*	Furcation involvement

Both the number and the * should be recorded if a furcation is detected - e.g. the score for a sextant could be 3* (e.g. indicating probing depth 3.5-5.5 mm PLUS furcation involvement in the sextant).

An example BPE score grid might look like:

4	3	3*
-	2	4*

When to record the BPE ?

- ▶ All new patients should have the BPE recorded
- ▶ For patients with codes 0, 1 or 2, the BPE should be recorded at least annually
- ▶ For patients with BPE codes of 3 or 4, more detailed periodontal charting is required:- Code 3: record full probing depths (6 sites per tooth) in the sextant(s) where the code 3 was recorded, in addition to recording the BPE in those sextants with scores 0, 1 or 2 . Code 4: if there is a code 4 in any sextant, then record full probing depths (6 sites per tooth) throughout the entire dentition
- ▶ BPE cannot be used to assess the response to periodontal therapy because it does not provide information about how sites within a sextant change after treatment. To assess the response to treatment, probing depths should be recorded at 6 sites per tooth pre- and post-treatment
- ▶ For patients who have undergone initial therapy for periodontitis (i.e. who had pretreatment BPE scores of 3 or 4), and who are now in the maintenance phase of care, then full probing depths throughout the entire dentition should be recorded at least annually.



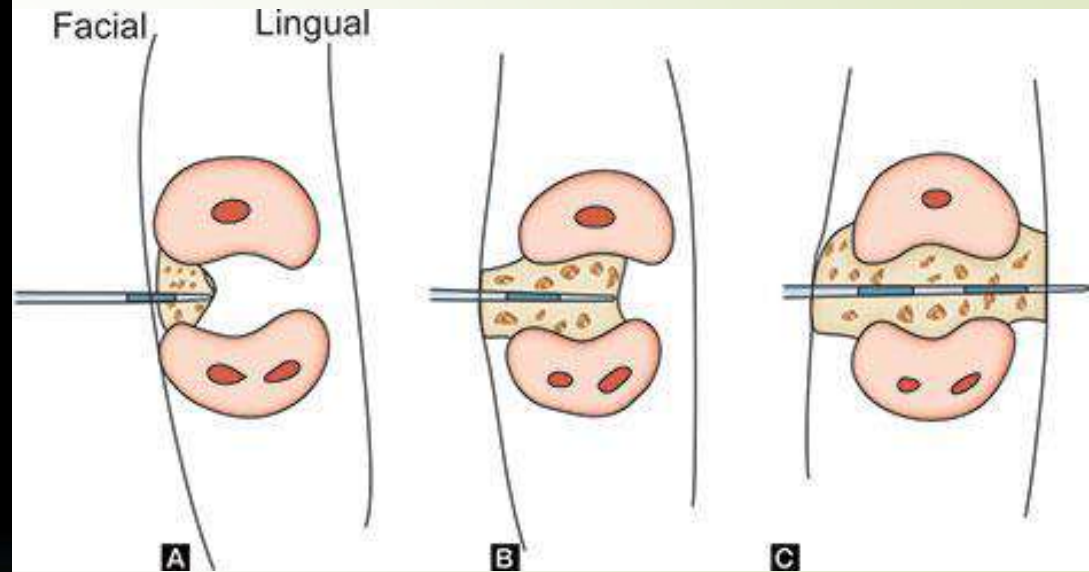
Guidance on interpretation of BPE score

0	No need for periodontal treatment
1	Oral hygiene instruction (OHI)
2	OHI, removal of plaque retentive factors, including all supra- and subgingival calculus
3	OHI, root surface debridement (RSD)
4	OHI, RSD. Assess the need for more complex treatment; referral to a specialist may be indicated.
*	OHI, RSD. Assess the need for more complex treatment; referral to a specialist may be indicated.

Furcation involvement index

Glickman's Classification(1953)

- Grade I Incipient Furcation
- Grade II cul-de-sac
- Grade III Communicating or Through and Through Furcation
- Grade IV Recession and clinically visible furcation



Recession index (Miller)

Table 1. Miller's classification of gingival recession defects.

	Symptoms	Treatment	Success
<i>Class I</i>	Recession that does not extend to the mucogingival junction	Complete root coverage is achievable	100%
<i>Class II</i>	Recession that extends to or beyond the mucogingival junction, with no periodontal attachment loss (i.e bone, soft tissue)	Complete root coverage is achievable	100%
<i>Class III</i>	Recession that extends to or beyond the mucogingival junction, with periodontal attachment loss in the interdental area or malpositioning of the teeth	Only partial root coverage possible to the height of the contour of interproximal tissue.	50-70%
<i>Class IV</i>	Recession that extends to or beyond the mucogingival junction, with severe bone or soft-tissue loss in the interdental area and/or severe malpositioning of the teeth	Root coverage is unpredictable and requires adjunctive treatment (ie orthodontics)	<10%

Recession index (Miller)





Thank you

بِسْمِ اللَّهِ الرَّحْمَنِ

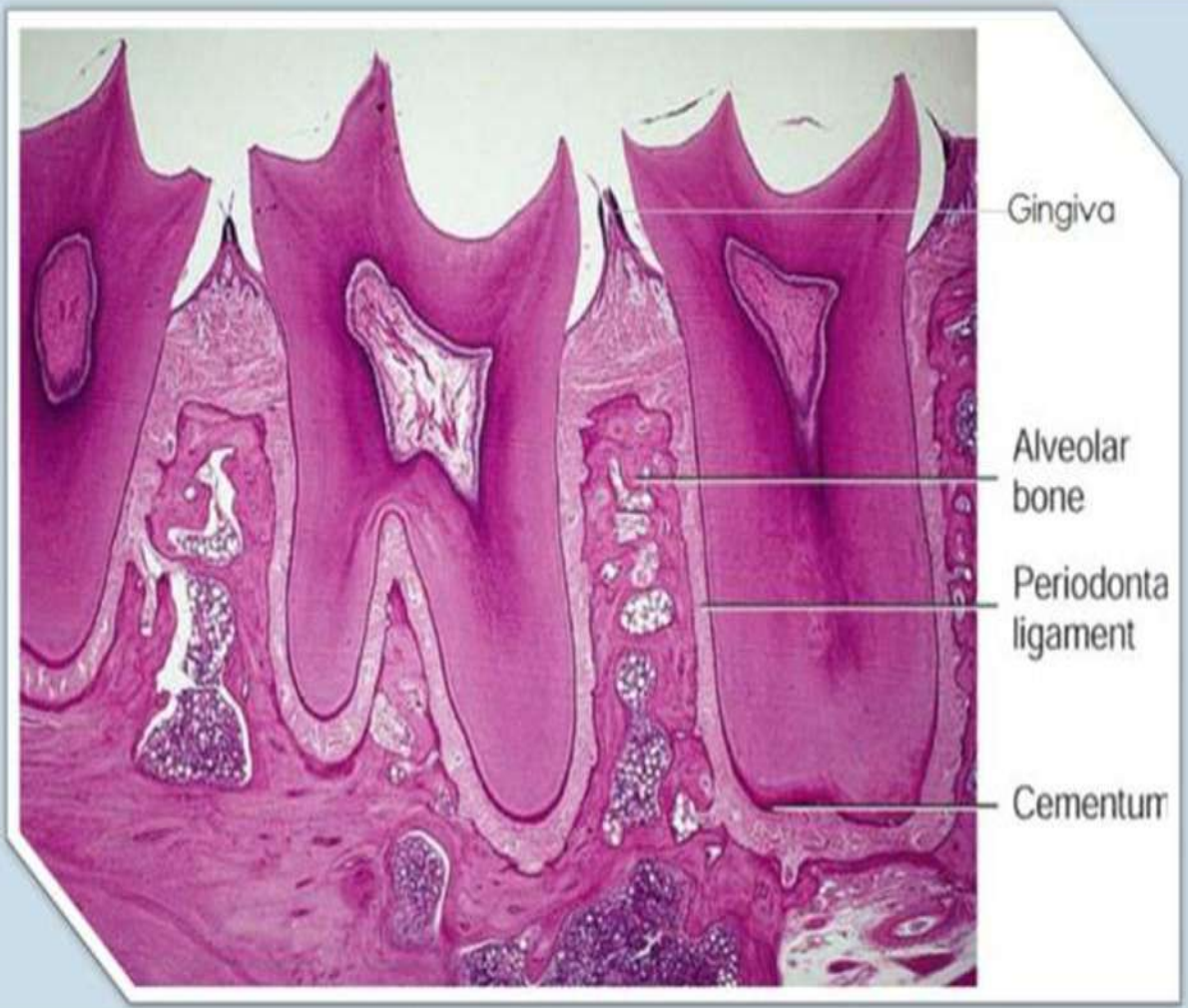
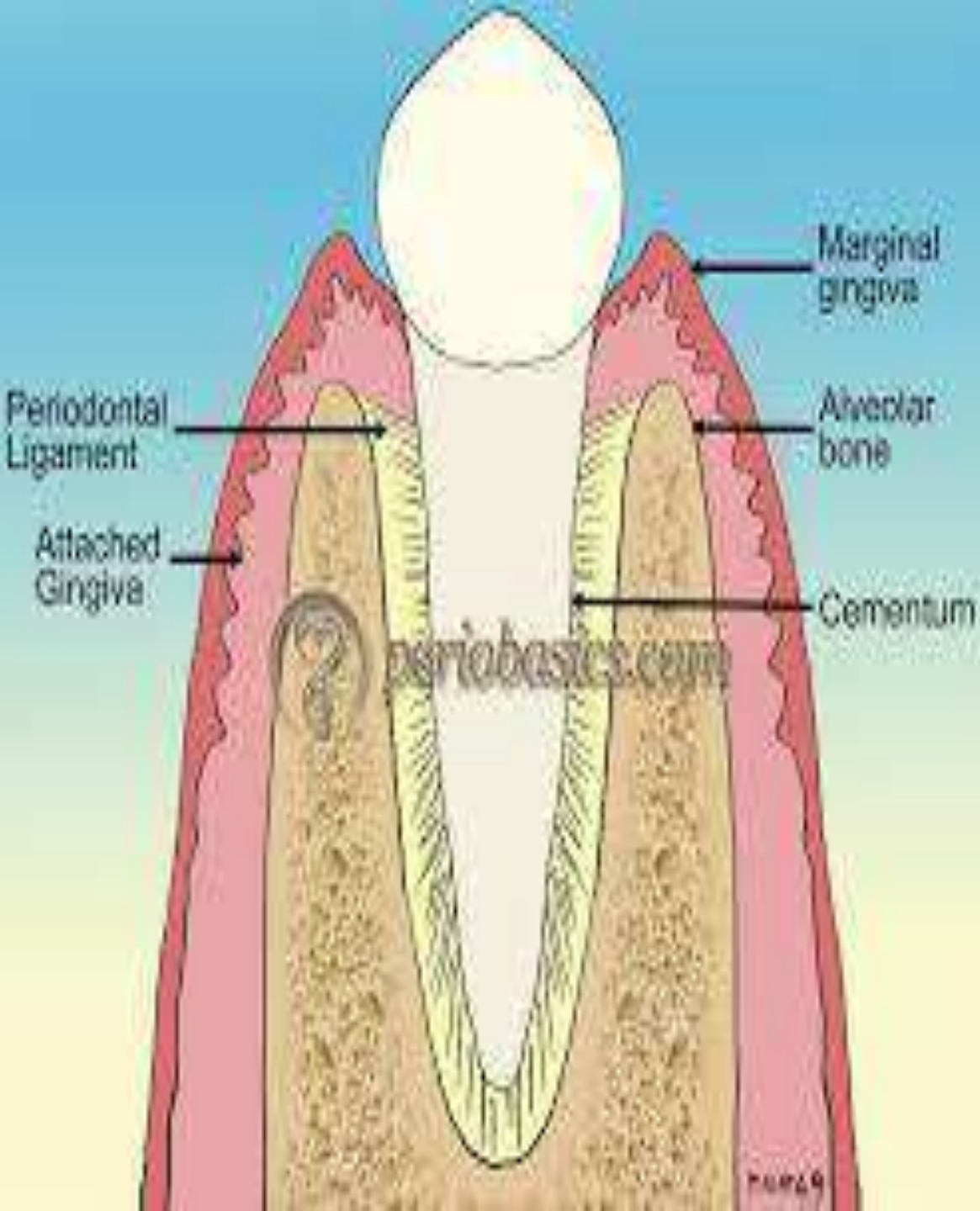
الرَّحِيمِ

PERIODONTAL LIGAMENT

أعداد:

د. نور صباح أرحيم

PERIODONTIUM



PERIODONTAL LIGAMENT (PDL)

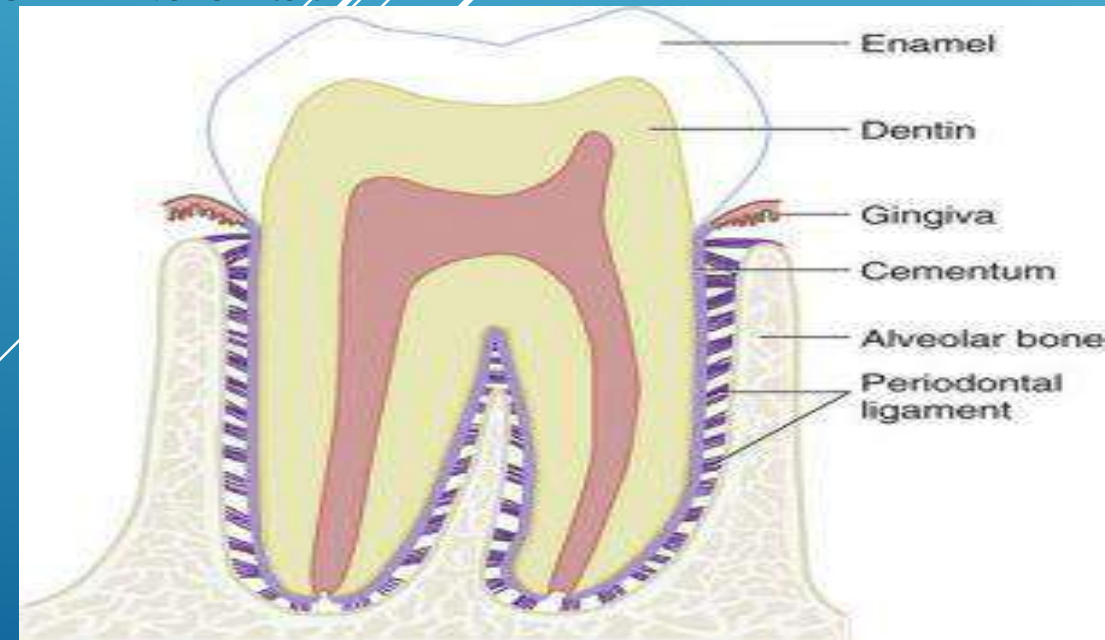
➔ It is a connective tissue surrounding the root and connecting it with the bone.

➔ It is consist primarily of:

1- Bundles of intermingling collagen fibers.

2- Cellular elements.

3- Ground substance.



DEVELOPMENT OF PDL

➔ The PDL and the cementum develop from follicular sac which derived from mesenchyme. The development of PDL occur during root formation and tooth eruption.

FIBERS OF PDL

((A)) The Majority of fibers in PDL are **collagens fibers**
(Principles fibers of PDL).

((B)) **Elastic fibers.**

((C)) **Oxytalan fibers.**

PDL

SHARPEY'S FIBERS:

It is the terminal portions of the principles fibers that insert into the cementum and bone.

Intermediate plexus:

It is the parts of the principle fibers, one is located toward the cementum and the other toward the alveolar bone and spliced together in the mid way between the cementum and bone to form intermediate plexus.

Sharpey's fibers

- The ends of the periodontal fibers that are embedded in alveolar bone and cementum are called Sharpey's fibers.
- On the cementum side these Sharpey's fibers are much thinner in diameter and insert at closer intervals as compared with the alveolar bone side

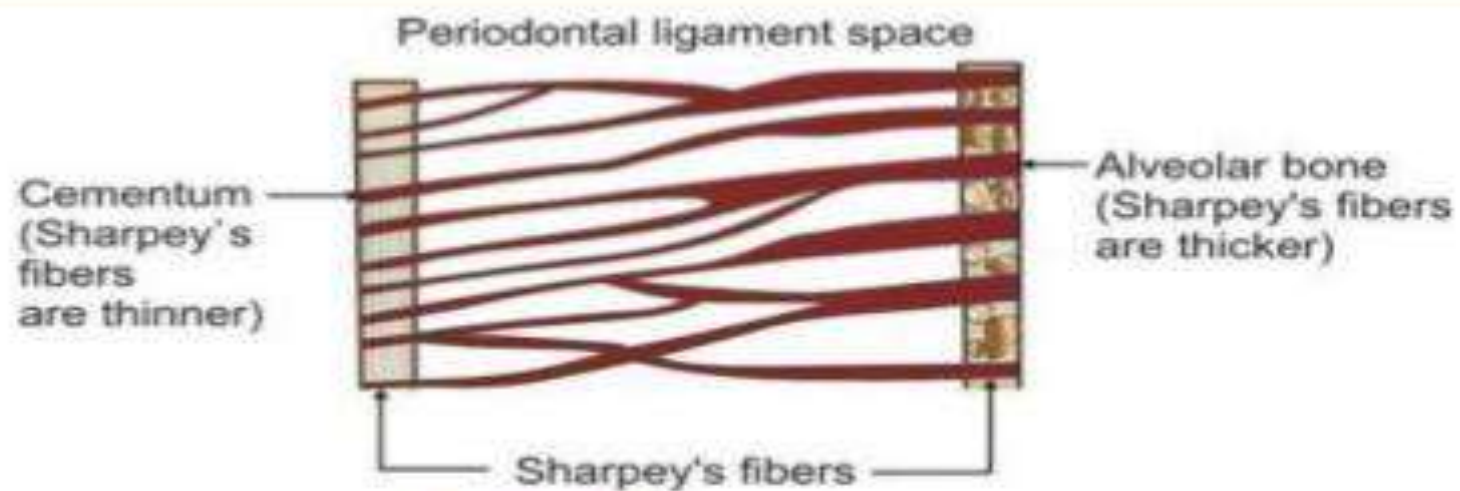
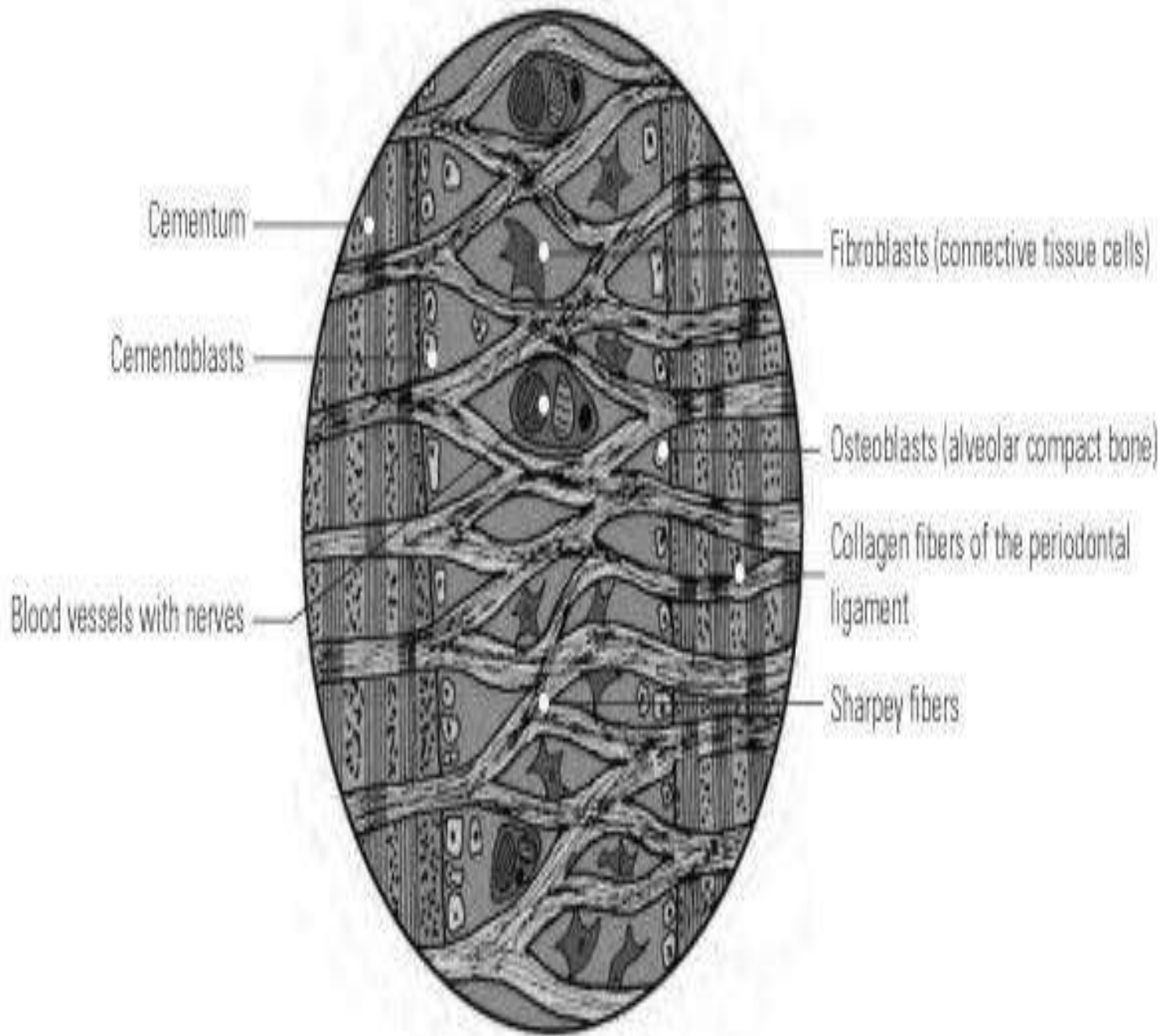
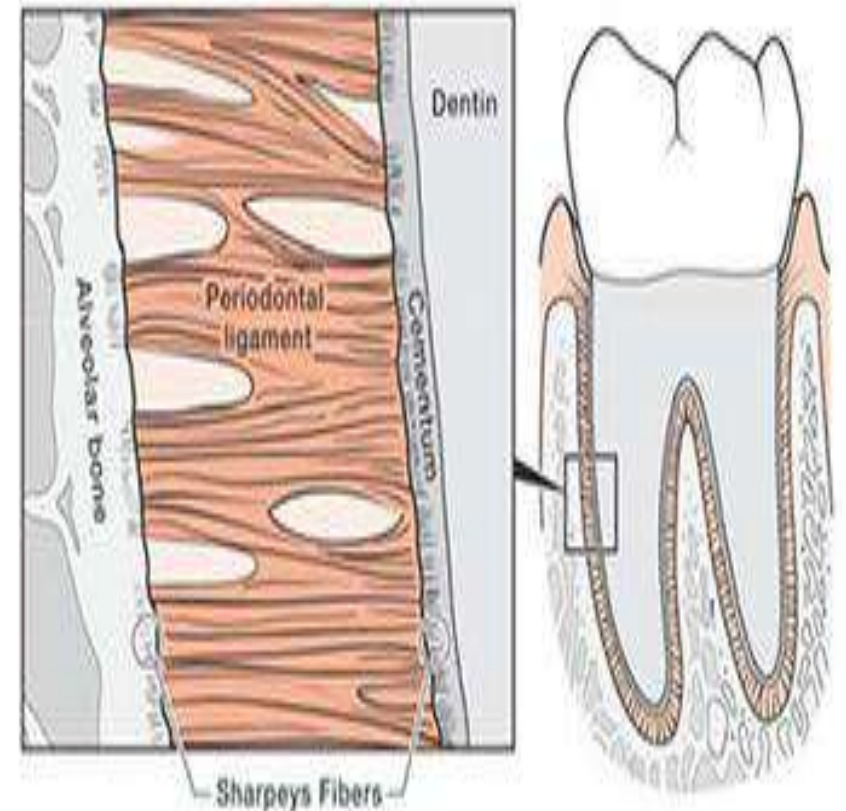


Fig. 2.3: Sharpey's fibers



Sharpey's Fibers



Copyright © 2008 Wolters Kluwer Health | Lippincott Williams & Wilkins

THE PRINCIPLES FIBERS OF THE PDL:

1- Alveolar crest fibers (ACP).

2- Horizontal fibers (HF).

3- Oblique fibers (OF).

4- Apical fibers (AF).

5- Interradicular fibers (IF).

6- Trasseptal Fibers (TF).



1-Alveolar crest fibers (ACF): ▶

They extend obliquely from the cementum to the crest of alveolar bone, they run in an apical direction. They prevent the extrusion of the tooth and resist lateral tooth movement. ▶

2-Horizontal fibers (HF): ▶

They extend at right angles to the long axis of the tooth from the cementum to the alveolar bone. ▶

3-Oblique fibers (OF): ▶

They are the largest group in the PDL, extend from the cementum in a coronal direction obliquely to the bone. They withstand the vertical masticatory forces. ▶

4-Apical groups (AP): ▶

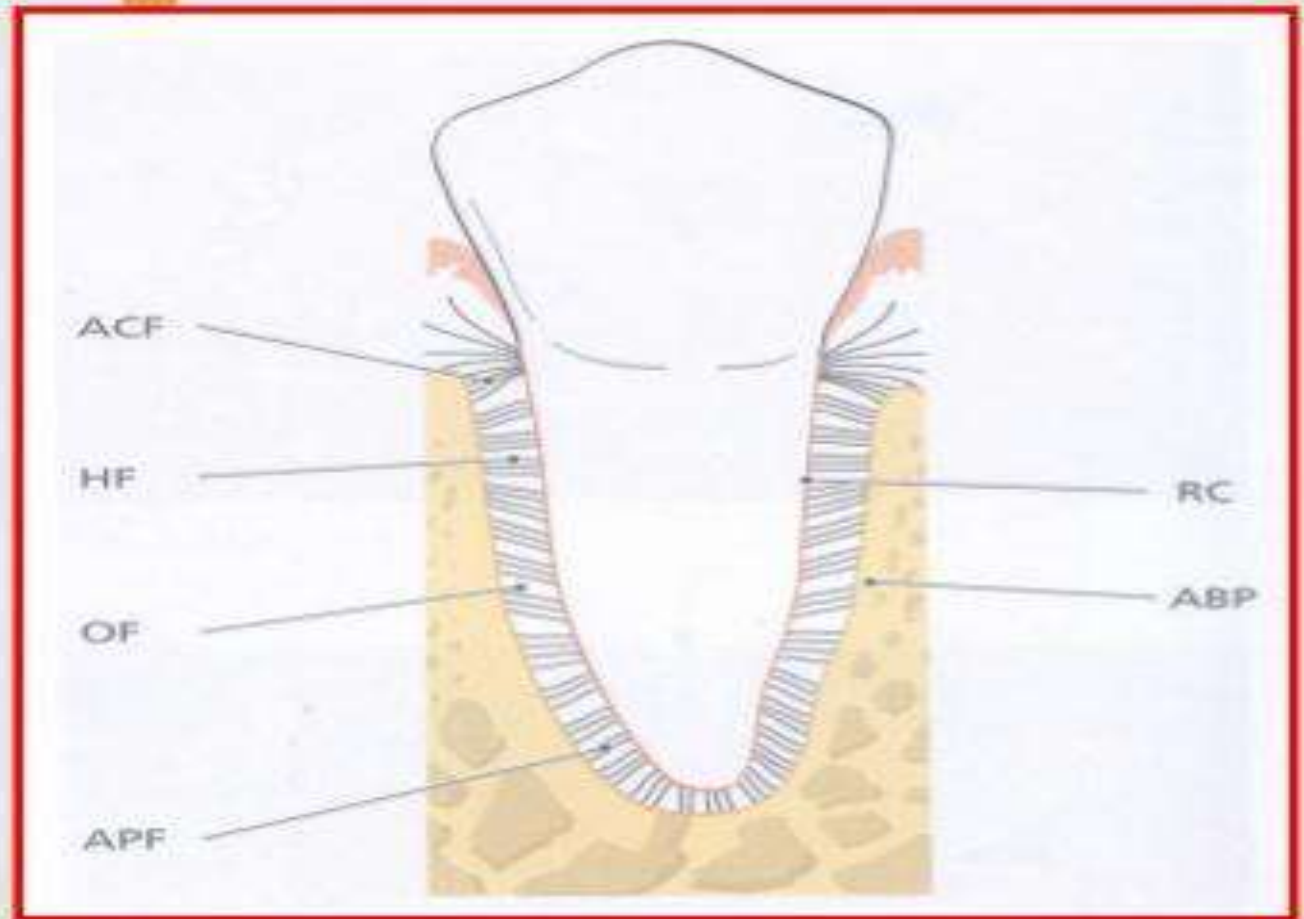
They radiate from the cementum to the bone at the apical region of the socket. ▶

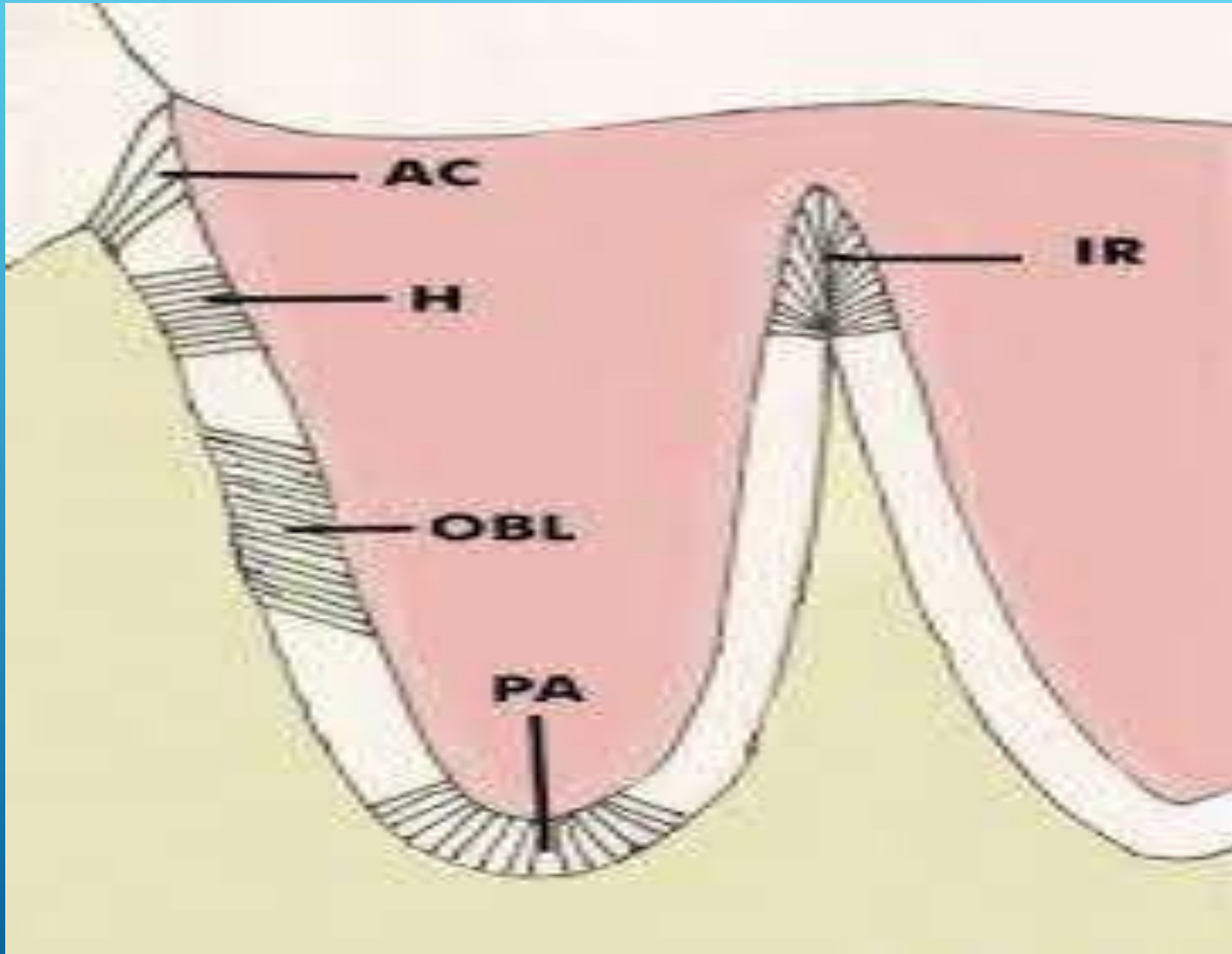
5-Interredicular fibers (IF): ▶

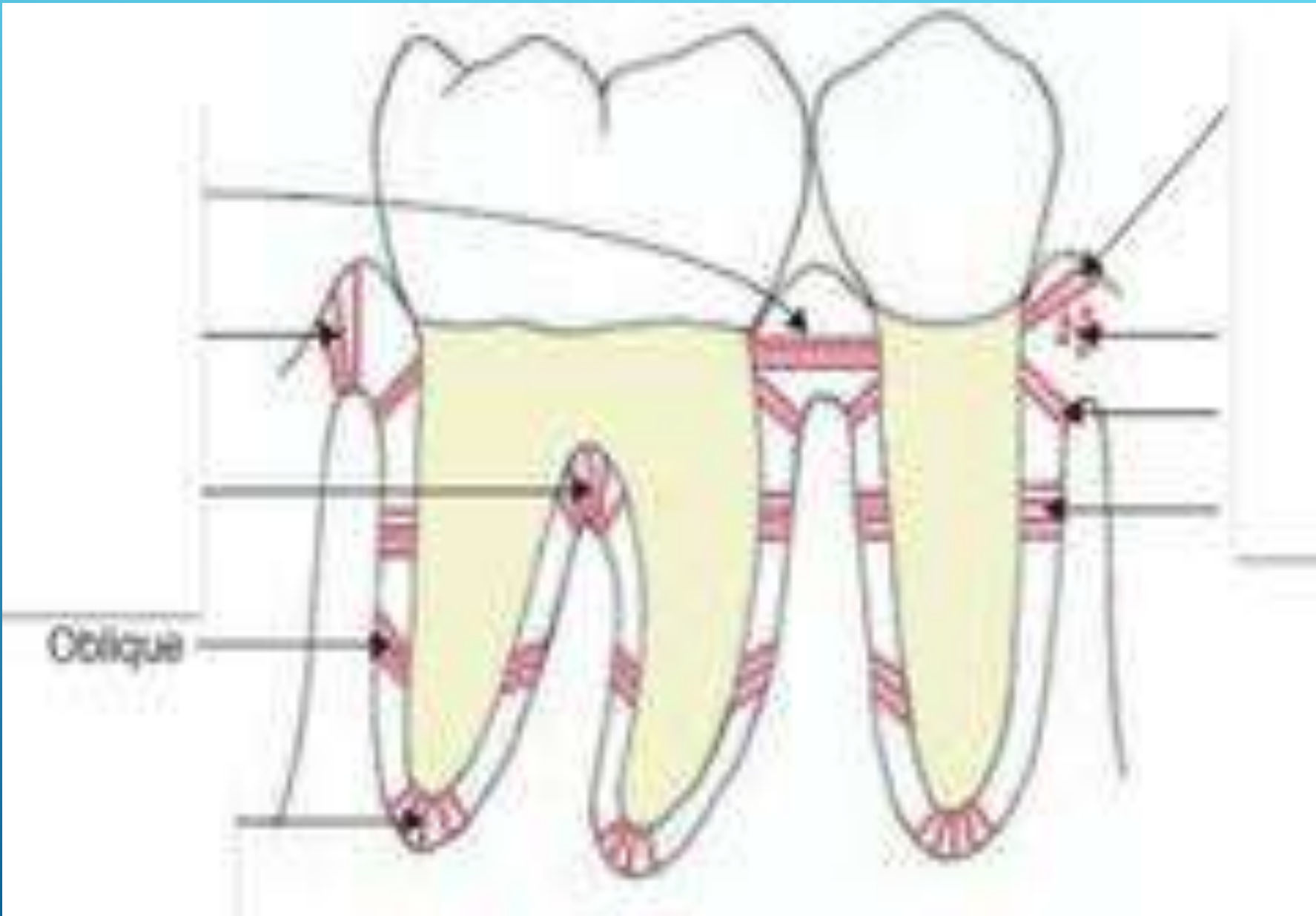
They run from the cementum to the bone in the furcation areas of multirooted teeth. ▶

Principal fibers of PDL

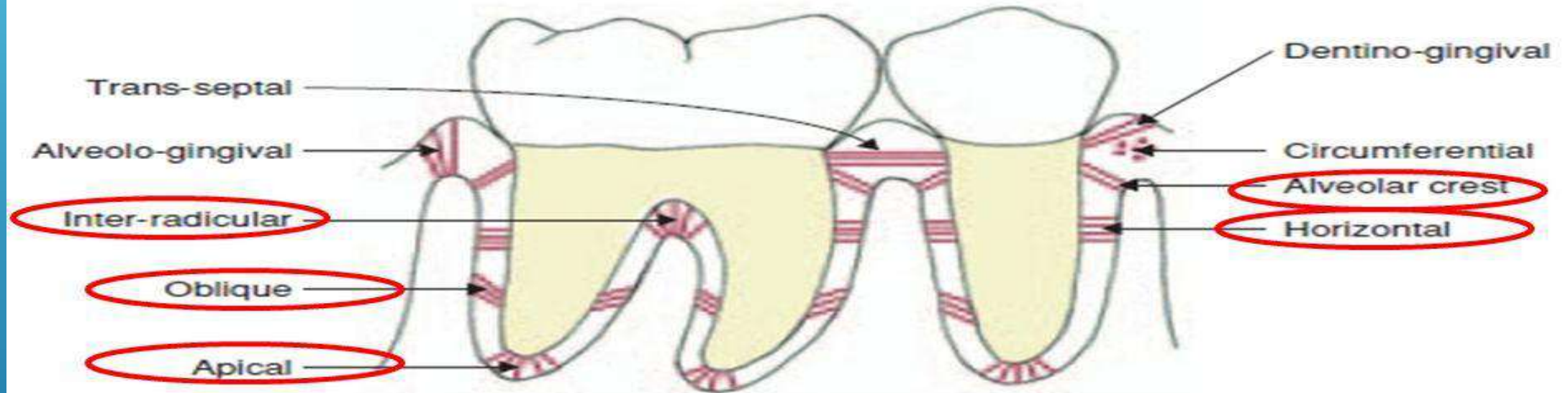
- Trans septal
- Alveolar crest group
- Horizontal
- Oblique
- Apical
- Inter - radicular







A- THE PRINCIPAL FIBERS



1- principle fiber bundle of the PDL:

- i. The alveolar crest group.
- ii. The horizontal group.
- iii. The oblique group.
- iv. The apical group.
- v. The interradicular group.

B-Elastic fibers: ▶▶

These are relatively few, and associated with the blood vessels. ▶

C- Oxytalan fibers: ▶

These immature forms of the fibers are thought to regulate vascular flow. ▶

CELLULAR ELEMENTS OF PDL:

1- Synthetic cells

(a) Osteoblasts

(b) Fibroblasts

(c) Cementoblasts

2- Resorptive cells

(a) Osteoclast

(b) Fibroblasts

(c) Cementoclasts

3- Epithelial rests of Malassez cells.

4- Immune system cells.

Osteoblasts: these cells cover the osseous surface of the PDL and are responsible for the formation of the alveolar bone. ▶

Fibroblast: these are the most common cells in the PDL. The main function of fibroblast is the production of various fibers such as collagen fiber, oxytalan fibers and elastic fibers. ▶

Cementoblasts: are seen lining the cementum surface of the PDL and are responsible for cementum deposition. ▶

Osteoclasts: these are large multinucleated cells and responsible for the bone resorption. ▶

Fibroblaste: these cells synthesize collagen and also possess the capacity to phagocytic old collagen fibers and degrade them by enzyme hydrolysis. The process of fibers resorption occur either during disease or physiological turnover. ▶

Cementoclast: cementum not remodeling as the alveolar bone and PDL but it undergoes continual deposition during life. ▶



Epithelial rest malassez:

these are found close to the cementum .
They are remnants of Hertwigs root sheath.
They proliferate when stimulated and
participate in the formation of periapical
cysts and lateral root cysts.

Immune system cells: ▶

Include mast cells, macrophage , lymphocyte ▶
neutrophil .

Decorative white lines consisting of several parallel diagonal strokes in the bottom right corner of the slide.

GROUND SUBSTANCE OF PDL:

A- Glycosaminglycan.

B- Glycoprotein.

➔ Water content 70%.



Principal fibers of the periodontal ligament

- | | |
|-----------------------|-----------------------------------|
| 1. transseptal fibers | 6. alveolar crest |
| 2. oblique | 7. dento-gingival (free gingival) |
| 3. apical | 8. alveolar-gingival |
| 4. interradicular | 9. circular |
| 5. horizontal | 10. dento-periosteal |

WIDTH OF PDL

- ➔ Width of PDL space is about 0.25mm.
- ➔ In hyperfunction, the tooth has wider PDL space.
- ➔ In hypofunction, the tooth has narrow PDL space.
- ➔ The space widest at the cervical and apical portion of the root and narrowest at the middle.
- ➔ in physiological tooth migration, the PDL space in the mesial root is thinner than the PDL space in distal root.
- ➔ PDL space decrease with age.
- ➔ in orthodontic treatment, the pressure side have smaller PDL space than tension side.
- ➔ in Implant and ankylois, there is no periodontal ligament space

Pdl space



Widened pdl space



Narrow PDL space



ELASTICITY OF PDL

- 1- Wavy coarse of the principal fibers.**
- 2- Intermediate plexuses.**
- 3- The Presence of oxytalan and Elastic fibers.**

FUNCTIONS OF THE PDL


1-Physical function.

2- Formative and Remodeling function.

3- Nutritive function.

4- Sensory function.

Physiological function: ▶


- 1-Transmission of occlusal forces to the bone.
 - 2-Attachment of the teeth to the bone.
 - 3-Resistance to the impact of occlusal forces(shock absorption).
 - 4-Provision of a soft tissue (casing) to protect the vessels and nerves from injury by mechanical forces.
- 
- A decorative graphic consisting of several parallel white lines of varying lengths and orientations, located in the bottom right corner of the slide.

Formative and remodeling function

Cells of the PDL participate in the formation and resorption of cementum and bone which occur in physiological tooth movement , in accommodation of the periodontium to occlusal forces and in the repair of injury.

Nutritive function

The PDL supplies nutrition to the cementum, bone and gingiva by way of the blood vessels and provide lymphatic drainage.

A decorative graphic consisting of several parallel white lines of varying lengths, slanted upwards from left to right, located in the bottom right corner of the slide.

Sensory function

The PDL is supplied with sensory nerve fibers which transmit tactile-pressure- pain sensation by the trigeminal pathway in addition the PDL is supplied with mechanoreceptor that transmit sense of localization.

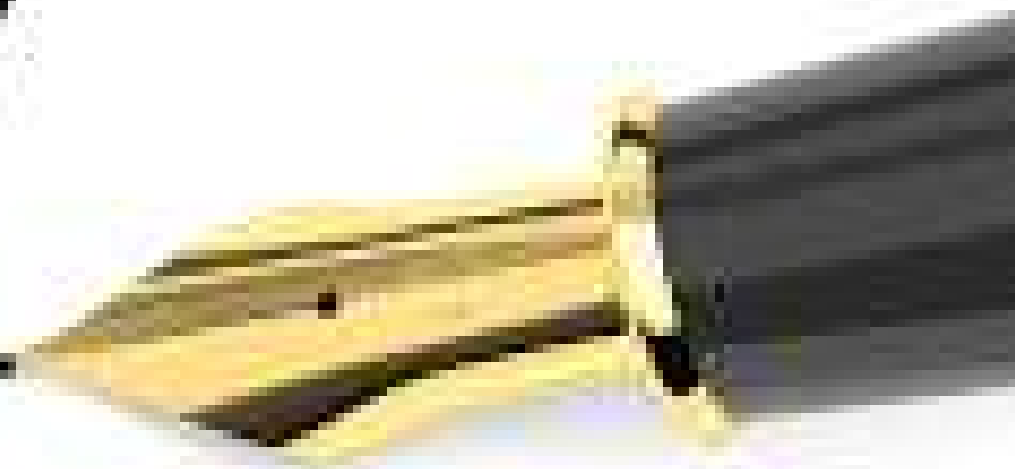
BLOOD SUPPLY OF PDL

● The blood supply to the supporting structures of the tooth is derived from the inferior and superior alveolar arteries to the mandible and maxilla respectively and reach the PDL from three sources:

- (A) Apical vessels supply the apical region of the PDL.**
- (B) The transalveolar vessels from the alveolar bone.**
- (C) Anastomosing vessels from the gingiva.**

Thank

you





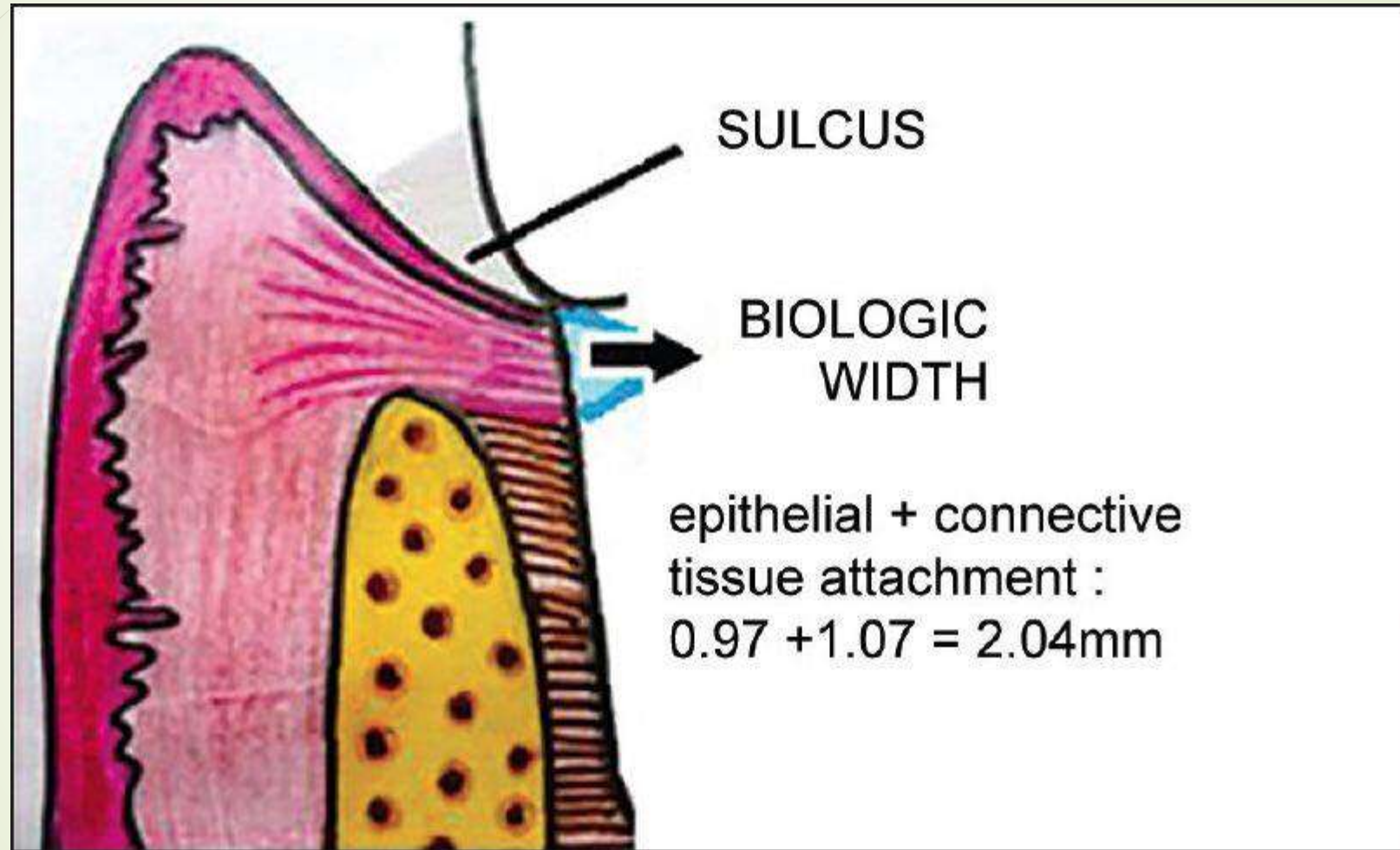
Gingival and periodontal pocket

م. سها أسود دهش العزاوي

B.D.S, MSC. PERIODONTOLOGY

Tooth gingival interface

- ▶ The interface between a tooth and the surrounding gingival tissue is a dynamic structure. The gingival tissue forms a crevice surrounding the tooth, resemble fluid-filled moat, The depth of this crevice, known as a sulcus, is in a constant state of flux due to microbial invasion and subsequent immune response. Located at the depth of the sulcus is the epithelial attachment, consisting of approximately 1 mm of junctional epithelium and another 1 mm of gingival fiber attachment, comprising the 2 mm of biologic width naturally found in the oral cavity.



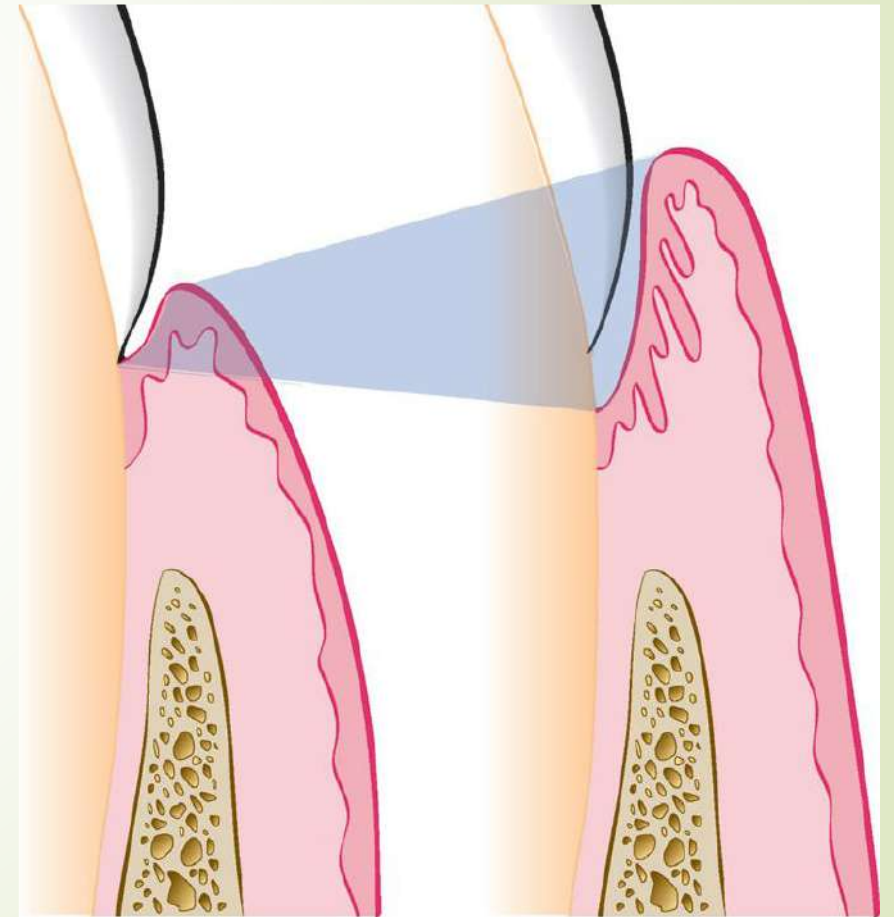
Gingival sulcus

A healthy sulcular depth is **3 millimeters or less**. Sulcular depths of 3 millimeters or less are readily self-cleansable with a properly used toothbrush or the supplemental use of other oral hygiene aids. When the sulcular depth is chronically in excess of three millimeters, regular home care may be insufficient to properly cleanse the full depth of the sulcus, allowing food debris and microbes to accumulate, forming dental biofilm. If accumulated microbes remain undisturbed in a sulcus for an extended period of time, they will penetrate and ultimately destroy the delicate soft tissue and periodontal attachment fibers. If left untreated, this process may lead to a deepening of the sulcus, recession, destruction of the periodontium, including the bony tooth socket, tooth mobility, and tooth loss.

➔ **Gingival and periodontal pockets** are dental terms indicating the presence of an abnormal depth of the gingival sulcus

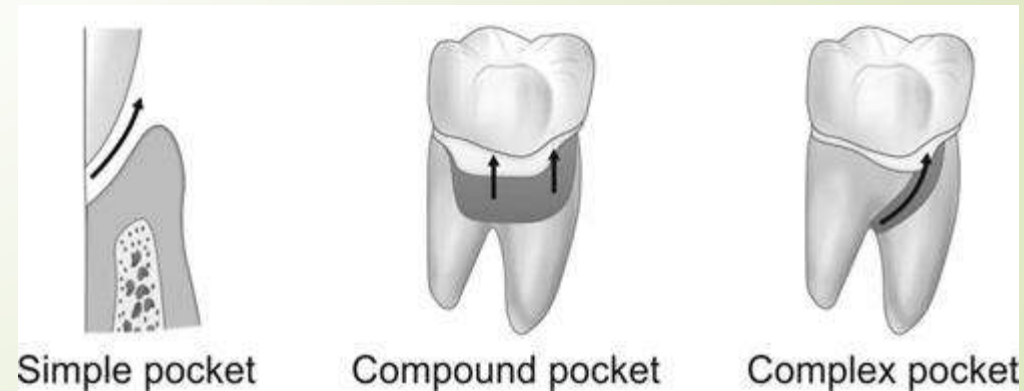
packet

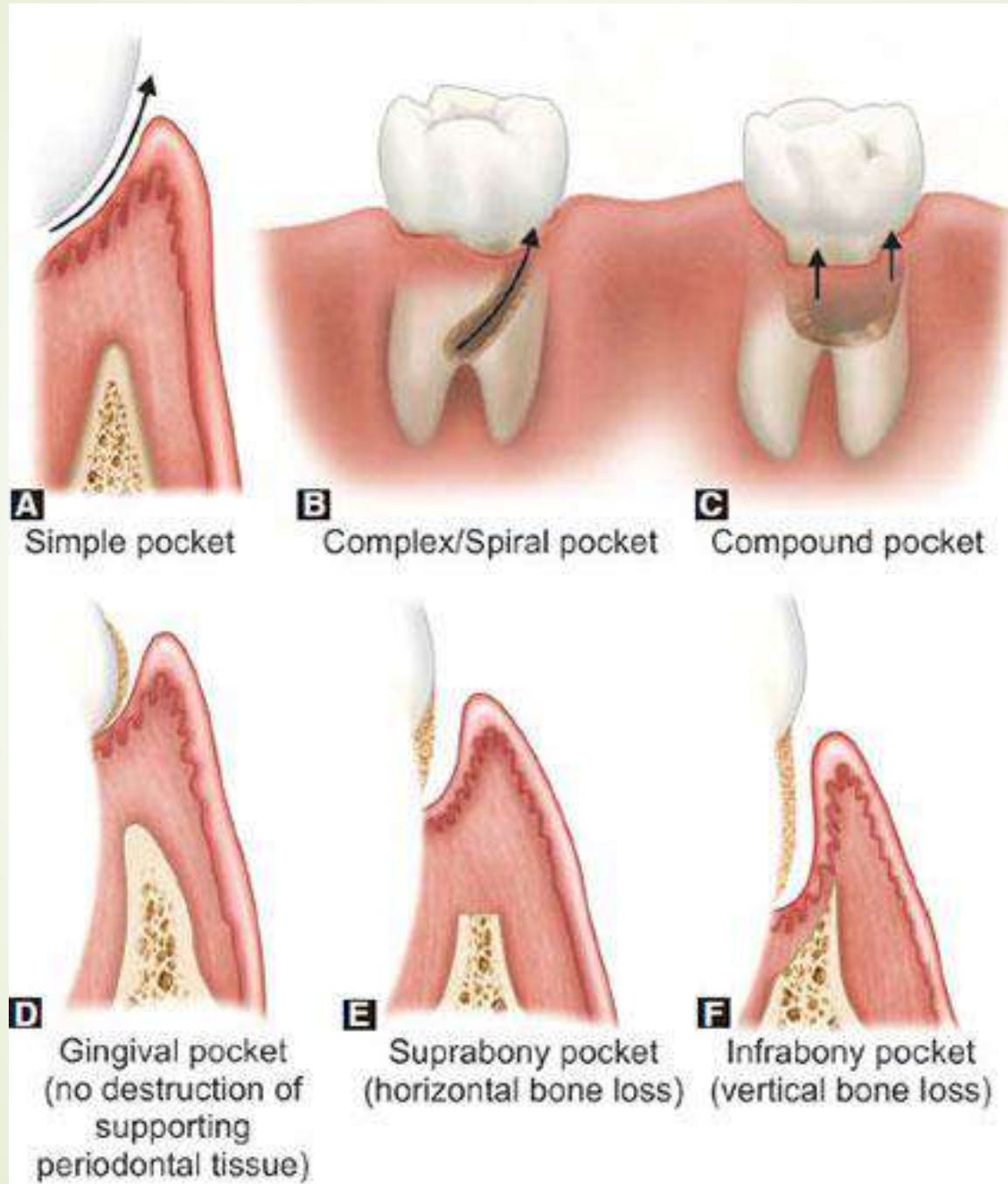
- It's an inflammatory changes and apical migration of junctional epithelium; it is also defined as a **pathological deepening of gingival sulcus**, which occurs by coronal movement of gingival margin, apical displacement of gingival attachment, or both.



Classification:

- **1-According to the involved tooth surface:**
 - Simple pocket: involve one surface
 - Compound pocket: involve more than one surface
 - Complex or spiral pocket: originating on one surface and twisting around the tooth to involve one or more additional surface. most common in furcation areas.


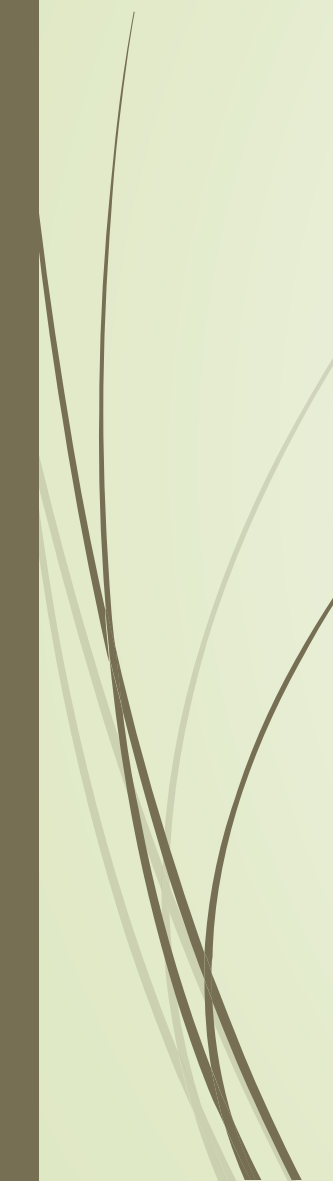




➤ 2-According to its location:

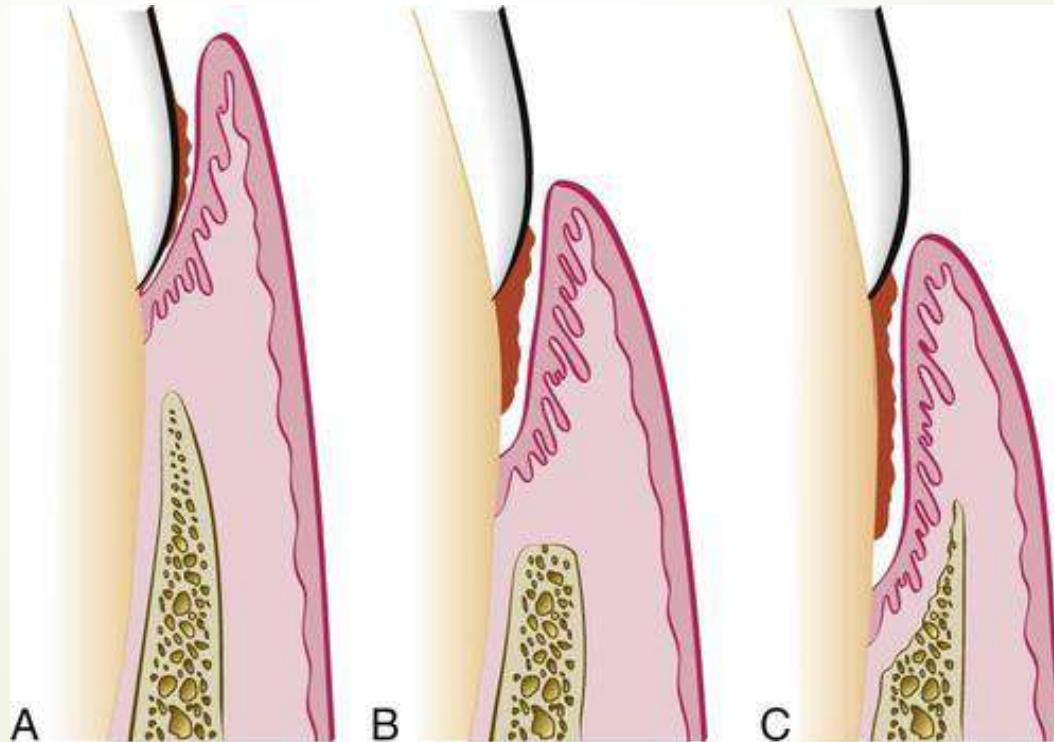
➤ **A- Gingival pocket:** which is formed by gingival enlargement without destruction of underlying periodontal tissue. The sulcus is deepened because of increased bulk of the gingiva.

- This phenomenon is also referred to as a false pocket or **pseudopocket**. The epithelial attachment does not migrate, it simply remains at the same attachment level found in pre-pathological health. The only anatomical landmark experiencing migration is the **gingival margin in a coronal direction**.
- In a gingival pocket, no destruction of the connective tissue fibers (gingival fibers) or alveolar bone occurs. This early sign of disease in the mouth is **completely reversible** when the etiology of the edematous reaction is eliminated and frequently occurs without dental surgical therapy.
- However, in certain situations, **gingivectomy** is necessary to reduce the gingival pocket depths to a healthy 1–3 mm.

- 
- **B- periodontal pocket**, which is defined as a pathologically deepened gingival sulcus, is one of the most important clinical features of periodontal disease. All different types of periodontitis, share histopathologic features, such as tissue changes in the periodontal pocket, mechanisms of tissue destruction, and healing mechanisms. However, they differ with regard to their etiology, natural history, progression, and response to therapy.
- 

3. According to its relation to alveolar crest:

- Suprabony pocket: also called supra crestal or supra alveolar. The base of the pocket is coronal to the level of underlying bone. The bone loss is horizontal
- Infrabony pocket: also known as sub crestal or intra alveolar pocket. The base of the pocket is apical to the level of adjacent bone. The bone loss is vertical.




Diagnosis/ detection of pockets

- Careful exploration with periodontal probe. (this method is accurate).
- Radiographic: pockets are not detected by the radiographic examination because pockets are soft tissue changes.
- A calibrated silver points or gutta percha points can be used with radiographic to assist in determining the level of attachment of periodontal pocket.



Pathogenesis

- ▶ The initial lesion in the development of periodontitis is the inflammation of the gingiva in response to a bacterial challenge. Changes involved in the transition from the normal gingival sulcus to the pathologic periodontal pocket are associated with different proportions of bacterial cells in dental plaque. Healthy gingiva is associated with few microorganisms, mostly coccoid cells and straight rods. Diseased gingiva is associated with increased numbers of Spirochetes and motile rods. However, the microbiota of diseased sites cannot be used as a predictor of future attachment or bone loss, because their presence alone is not sufficient for disease to start or progress.



Sequences in pathogenesis of periodontal pocket

- Accumulation of microorganisms on the supra gingival tooth surface and its extension into gingival sulcus.
- Inflammatory changes in the connective tissue wall of the gingival sulcus.
- Cellular and fluid inflammatory exudate causes degeneration of the connective tissue including the gingival fibers.
- Collagen fibers gets destroyed apical to the junctional epithelium and the area becomes occupied by the inflammatory cells and edema.
- The coronal portion of the junctional epithelium detaches from the root as the apical portion migrates.
- Polymorphonuclear neutrophils invade the coronal end of the junctional epithelium in increasing number.
- With continued inflammation the gingiva increase in bulk and the crest of the gingival margin extends coronally.
- The junctional epithelium continues to migrate along the root and separate from the root.

Mechanisms of collagen loss:

Two mechanisms involved:

➤ First mechanism:

Collagenases and other enzymes secreted by fibroblast, PMNs and macrophages in healthy and inflamed tissues, become extracellular and destroy collagen.

➤ Second mechanism:

Fibroblast phagocytize collagen fibers by extending cytoplasmic processes to the ligament-cementum interface and degrade the inserted collagen fibrils and the fibrils of cementum matrix.




Content of the pocket:

- 1) Microorganisms .
- 2) Bacterial products (enzymes and endotoxins).
- 3) GCF.
- 4) Remnants of food
- 5) Salivary mucin.
- 6) Desquamated epithelial cells.
- 7) Leukocytes
- 8) Plaque covered calculus usually projects from the tooth surface
- 9) Purulent exudates may be present

Periodontal pocket as healing lesions:

- ▶ Periodontal pockets are inflammatory lesions and constantly undergoing repair. Complete healing does not occur because of persistence of bacterial attack which continues to stimulate an inflammatory response causing degeneration of new tissues elements formed during the continuous effort at repair.
- ▶ The condition of the soft-tissue wall of the periodontal pocket results from the interplay of the destructive and constructive tissue changes. Their balance determines clinical features such as color, consistency, and surface texture of the pocket wall..

- 
- ➔ **Edematous pocket walls:** when the **inflammatory component predominates** (inflammatory fluid and cellular exudate predominate) the lateral wall appears soft, edematous friable, with smooth shiny surface and bluish red discoloration.
 - ➔ **Fibrotic pocket wall:** when **reparative changes predominate**, (If there is a relative predominance of newly formed connective tissue cells and fibers), the pocket wall is more firm and pink the gingiva appears fibrotic and pink.
 - ➔ In some cases both lesions present in the same pocket as outer surface of a pocket wall fibrotic, the inner surface of soft tissue wall is inflamed and ulcerated.

Clinical features/histopathological feature:

A.1- bluish red discoloration of the gingival wall of pocket, this caused due to circulatory stagnation.

2- flaccidity of tissue: due to destruction of gingival fibers.

3- smooth shiny surface: due to atrophy of the epithelium and edema.,

4- pitting on pressure: due to edema and degeneration.

B. Gingival wall may be pink or firm when fibrotic changes predominates over exudation and regeneration.



C. Bleeding on probing: due to
Increased vascularity.

Thinning and degeneration of epithelium.

Proximity of engorged vessels to inner surface.

D. Probing is generally painful: due to ulceration of the inner
aspect of the pocket wall.

E. Pus may be present: due to suppurative inflammation.

F. Other clinical features

Thickened marginal gingiva.

Loss of stippling.

Tooth mobility and diastema formation

Periodontal disease activity

1- **Period of quiescence or inactivity** this period characterized by reduced inflammatory response and little or no loss of bone and connective tissues.

A buildup of unattached plaque with its gram negative and anaerobic bacteria

2- **period of exacerbation or activity** bone and connective tissue attachment are lost and the pocket deepens

This period may last for days, weeks, months and eventually followed by period of remission and quiescence in which G+ve bacteria proliferate and more stable condition is established

Clinical feature shows bleeding spontaneous or on probing and greater amount of gingival exudates

Histological features , pocket appear thin and ulcerated ,infiltrate composed of plasma cells and PMNs leukocytes



Pocket probing:

We have two different pocket depths:

- **Biologic or histologic depth:** distance between gingival margin and base of the pocket. measured histologically (accurate measurement but not used routinely)
- **Clinical or probing depth:** distance to which a probe penetrates into the pocket.
- The standardized force used for penetration of probe is 25 pounds or 23 grams (0.75 N).

Probing Pocket depth PPD: Distance between base of pocket and gingival margin.

➤ **Extent**

➤ The "extent" of disease refers to the proportion of the dentition affected by the disease in terms of percentage of sites. Sites are defined as the positions at which probing measurements are taken around each tooth and, generally, six probing sites around each tooth are recorded, as follows:

➤ Mesiobuccal

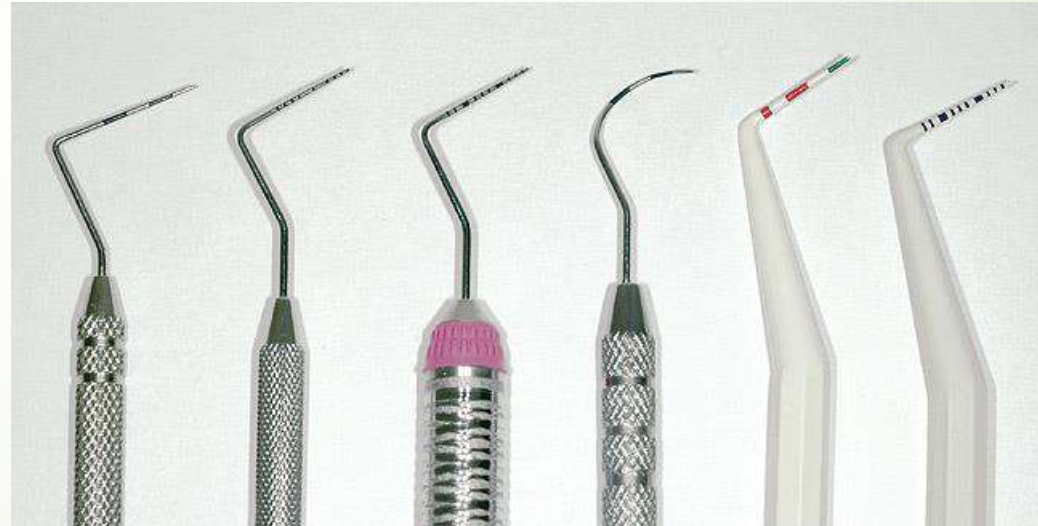
➤ Midbuccal

➤ distobuccal

➤ mesiolingual

➤ mid-lingual

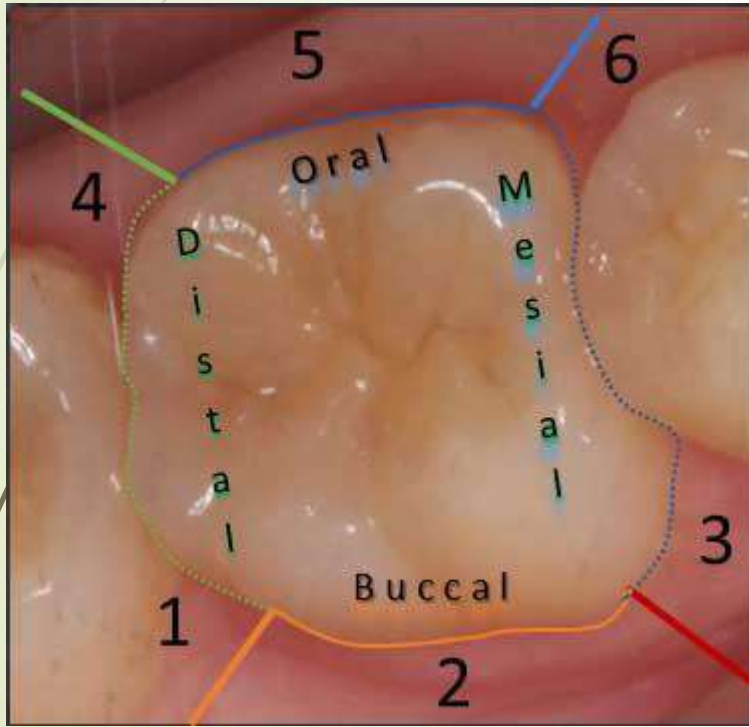
➤ distolingual



➤ If up to 30% of sites in the mouth are affected, the manifestation is classified as "localized"; for more than 30%, the term "generalized" is used.

Probing techniques

- Occlusal view: six surfaces measured in periodontal probing
- In multirrooted teeth, the possibility of furcation involvement should be carefully explored with specially designed probe (eg. **Nabers probe**).
- The probe should be inserted **parallel** to the vertical axis of the tooth and **walked** circumferentially around each tooth to detect the area of **deepest** penetration.
- To detect **internal crater**: the probe should be placed **obliquely** from both facial and lingual surfaces, so as to explore the deepest point of the pocket located beneath the contact point.



Level of attachment loss CAL

Distance between base of pocket and a fixed point on the tooth such as CEJ.

Severity: The "severity" of disease refers to the amount of periodontal ligament fibers that have been lost, termed "clinical attachment loss"

- STAGE 1: 1-2mm *4-5mm pocket*
- STAGE 2: 3-4mm *6-7mm*
- STAGE 3: >5mm *>8mm*
- STAGE 4: >5mm *>8mm*

Treatment:

1-Non surgical treatment:

- ▶ Oral hygiene instruction.
- ▶ Scaling and root planning
 - Use curettes for subgingival scaling, root planing and removal of the soft tissue lining the pocket.
 - Root planing stroke should be moderate to light.
 - Pull stroke for final smoothing and planing of root surface.
 - Continuous series of long, overlapping shaving stroke is achieved.
 - To avoid over instrumentation, a delicate transition from short, powerful scaling strokes to longer, lighter root planning strokes must be made as soon as calculus and initial roughness have been eliminated.
 - Periodontal medication as application of tetracyclines



2-Surgical treatment

Pocket depth reduction through different surgical procedures

- 1-gingival curettage
- 2-gingivectomy
- 3-periodontal flap procedures
- 4-osseous surgery
- 5-Periodontal regeneration procedures



Thank you

Phase II Surgical Therapy

م. سها أسود دهش العزاوي

B.D.S, MSc. Periodontology

Emergency Phase

Non-Surgical Phase

Maintenance Phase

Surgical Phase

Restorative Phase



Reevaluation after phase 1 therapy

All patients should be treated initially with scaling and root planing and the final decision on the need for periodontal surgery should be made only after evaluation of the effects of phase 1 therapy no less than 1 to 3 months after completion of phase one therapy.

- Patients with adequate oral hygiene and no remaining pockets: put in maintenance phase.
- Patients with inadequate oral hygiene: repeat phase 1 focusing on OHI.
- Patients with adequate oral hygiene but remaining pockets of $\geq 5\text{mm}$: periodontal surgery for pocket elimination.

In this phase the surgical techniques used for the following purposes :

- 1-Controlling or eliminating periodontal disease(surgical pocket therapy)
- 2-Correcting anatomic conditions that favor periodontal disease, impair aesthetics or impede placement of prosthetic appliances(plastic surgery, aesthetic surgery, pre prosthetic techniques).
- 3-Placing implants to replace lost teeth and improving environment for their placement and function.

Periodontal surgery

1. Pocket reduction surgery:

- **Resective** (gingivectomy, apically displaced flap and un-displaced flap with or without osseous resection).
- **Regenerative** (flaps with grafts, membranes, etc).

2. Correction of anatomic/morphologic defects:

- **Esthetic surgery** :e.g:covering deuded roots and augmentation of gingiva.
- **Preprosthetic surgery** (crown lengthening, ridge augmentation and vestibular deepening)

3. Placement of dental implants :

Periodontal surgery

- Successful cause-related therapy (by the removal of plaque and calculus) will reduce gingival inflammation (edema, hyperemia and flabby tissue) there by making assessment of true gingival contour and pocket depth possible. In addition the soft tissue will be more fibrous and thus firmer, which facilitate surgical handling of the soft tissues. The propensity for bleeding is reduced, making the inspection of the surgical field easier.

Periodontal surgery

- The effectiveness of the patient's home care which is of decisive importance for the long term prognosis must be properly evaluated; lack of effective self performed plaque control will often mean that the patient should be excluded from surgical treatment.

Objectives of periodontal surgery

- 1-Accessibility and direct vision for proper S+ RP.
- 2-Reduction or elimination of plaque retentive area especially periodontal pockets that have not responded to initial therapy.
- 3-Eliminate inflamed periodontal tissue.
- 4-Create a physiologic morphology of the dentogingival area that will facilitate efficient self performed plaque control.

-
- 5-Provide access to correct bony defects.
 - 6-Enhancing the regeneration of periodontal tissue.
 - 7-Correct mucogingival defect and improve periodontal aesthetic.

Surgical treatment include

- 1-Gingivectomy.
- 2-Flap surgery.
- 3-Distal wedge procedure.
- 4-Mucogingival surgery for correction of mucogingival and aesthetic defect.
- 5-Crown lengthening to increase clinical crown length
- 6-Guided tissue regeneration (GTR) to regenerate periodontal supporting structures.

Gingivectomy

- This surgical procedure aimed at the excision of the soft tissue wall of a pathologic periodontal pocket and this pocket elimination was usually combined with recontouring of the diseased gingiva to restore physiologic form(e.g. Drugs induced gingival enlargement and the resulting false pocket can be removed by this method).

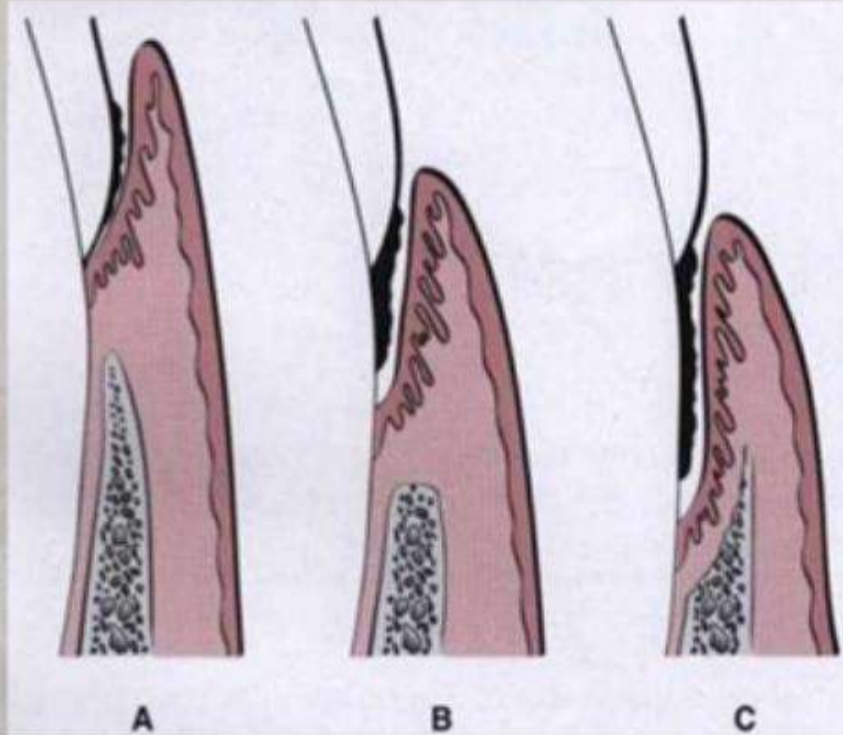
Indication

- 1-Gingival enlargement or over growth.
- 2-Idiopathic gingival fibromatosis.
- 3-Shallow suprabony pocket.
- 4-Minor corrective procedure.

1. According to the relation to the
crestal bone

Suprabony
supracrestal/
supraalveolar

Intrabony
subcrestal/
intraalveolar



Contraindication

- 1-Infrabony pocket.
- 2-Thickening of marginal alveolar bone and the need for bone surgery.
- 3-Attached gingiva is narrow or absent.

Advantages

- 1-Technically simple, good visual access.
- 2-Complete pocket elimination.
- 3-Restoration of a physiologic gingival contour.

Disadvantages

- 1-Gross wound, post operative pain.
- 2-Healing by secondary intention.
- 3-Danger of exposing bone.
- 4-Loss of attached gingiva.
- 5-Phonetics and aesthetic problem in the anterior area with sensitivity due to exposure of the cervical area of tooth.

1. SURGICAL GINGIVECTOMY

Instruments Required In Surgical Gingivectomy

- Krane Keplan Pocket Marker
- Kirkland Periodontal knife
- Orban periodontal knife
- Bard – parker handle
- Bard – Parker blades no 11 & 12
- Supra & subgingival scalers
- Curettes

Pocket marking forceps



- Paired (L & R)
- **Use:** indicate the location of the base of

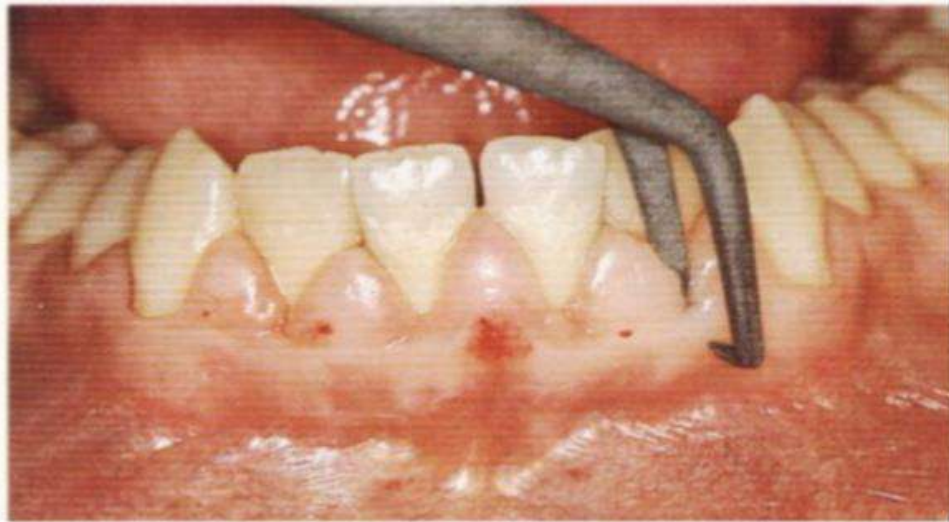
Gingivectomy knives



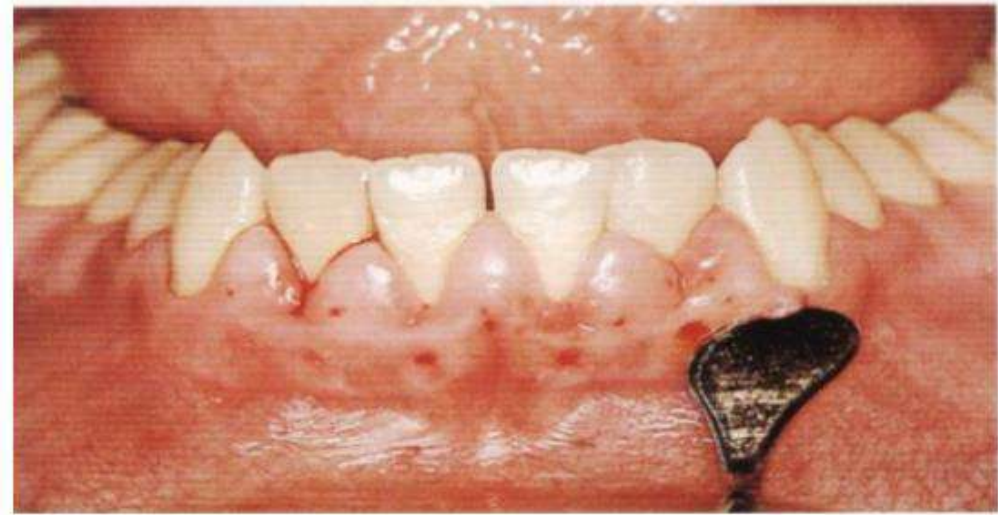
- GV knife (Kirkland, L & R)
- Papilla knife (Orban, L & R)
- Universal knife

Gingivectomy procedure

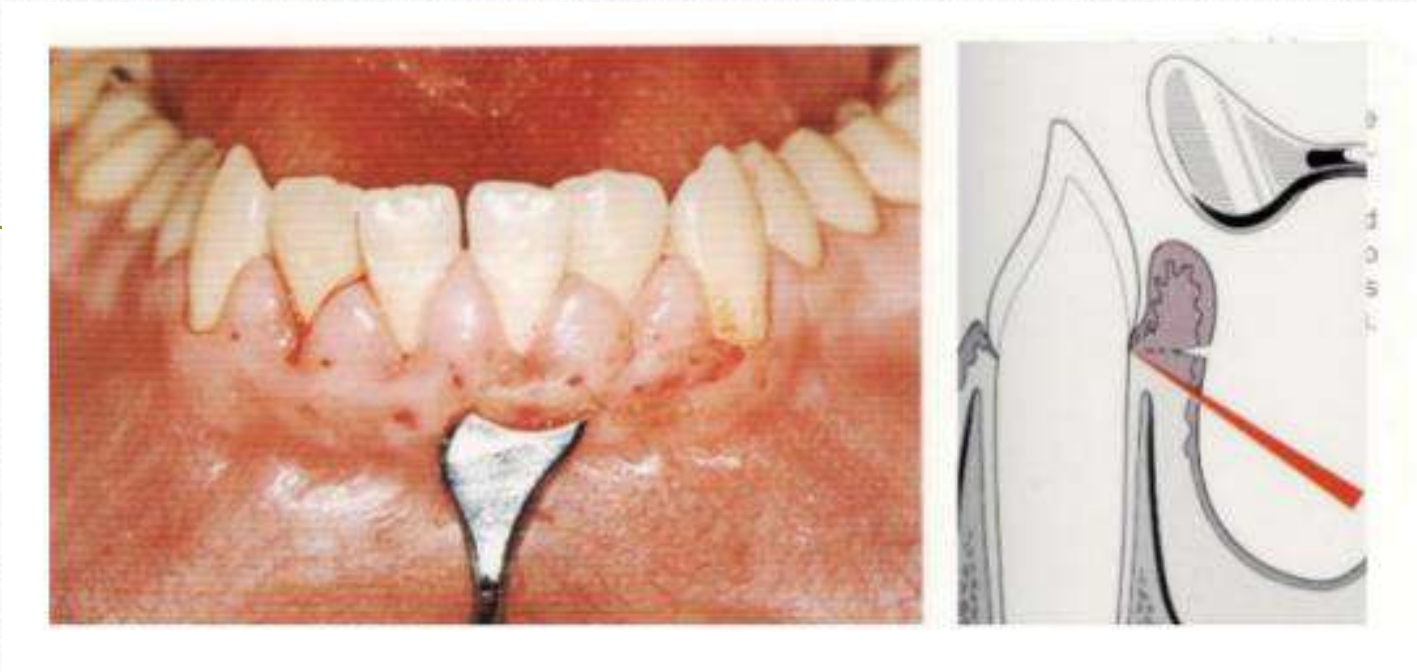
- Giving local anesthesia.
- Marking the pocket depth: the straight arm of pocket depth marker forceps is guided into buccal pocket, when the base of pocket is encountered, the forceps is pinched together causing the horizontal forceps tip to mark depth of pocket, by repeating this procedure at each tooth surface, a series of bleeding points is created, which are used subsequently as a guide for incision.



Marking the base of the pocket



Primary beveled incision which carried out slightly apical to bleeding points by Kirkland knife.



Continuous incision or interrupted , straight or scalloped is made.



Secondary incision to separate the interproximal soft tissues from the interdental periodontium by Orban knife.



Careful removal of the incised tissues by a curette or a cumine.



By curette remove plaque, calculus and granulation tissues then smoothing teeth surfaces.



Gingivectomy wound after scaling



Use Kirkland knife for gingivoplasty (minor alterations in gingival morphology without tissue excision) by shaving wound margin to create thin margin.



Control bleeding by placing gauze packs Put dressing to cover the wound with pressure to prevent the bleeding with consequence formation of granulation tissue under dressing and without interference with occlusion or mobile mucosa

Flap surgery

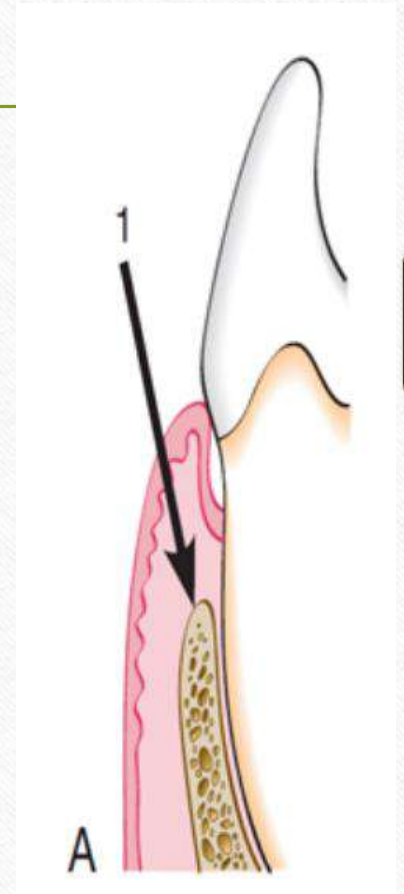
- **Indications**
- 1-In treatment of infrabony pockets.
- 2-When the gingivectomy will lead to an unacceptable aesthetic results.
- 3- Osseous recontouring (elimination of bony defect).

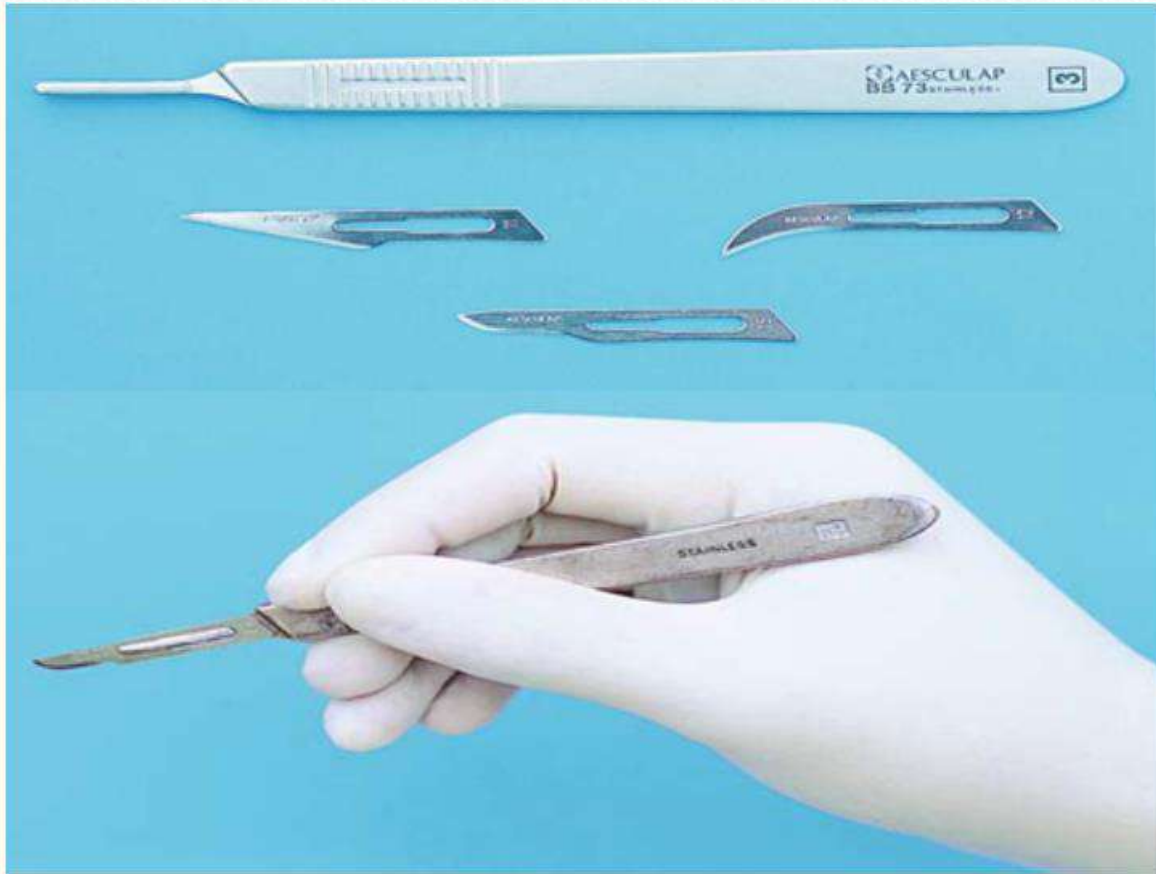
Modified widman flap

- Reported in 1974 by Ramfjord and Nissle, it is a replaced flap. There are three incisions in this flap, it is usually conducted as following:

A- Primary incision: First incision-scalloping

- The scalloped incision is performed on both labial and palatal aspects, using the double-edge 12B scalpel. It is an inverse bevel incision extending to the alveolar crest. This incision thins the gingival tissue and permits complete closure of the interdental osseous defects postoperatively. The distance of the incision from the gingival margin may vary from 0.5 to 2mm.





B- Flap retraction:

- An elevator is used to raise a full thickness mucoperiosteal flap as atraumatically as possible. The flap is reflected only to permit direct visualization of the root surface and the alveolar crest. In most cases it is possible to stay within the boundaries of the attached gingiva, without extending beyond the mucogingival line.



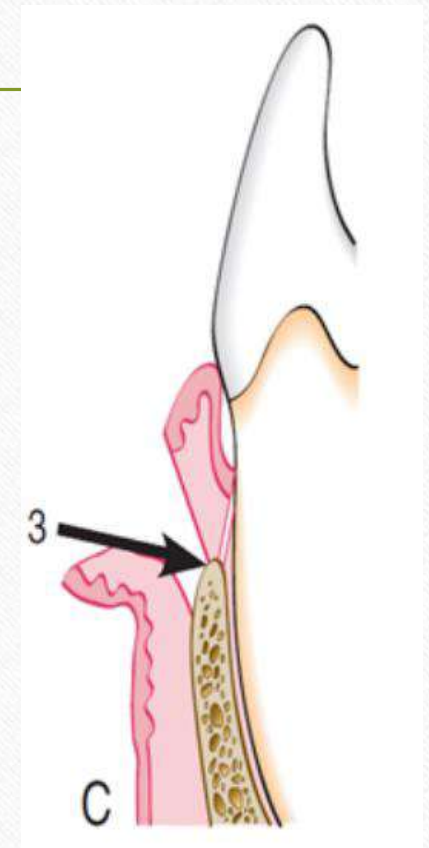
C-Secondary incision: Second incision-crevicular

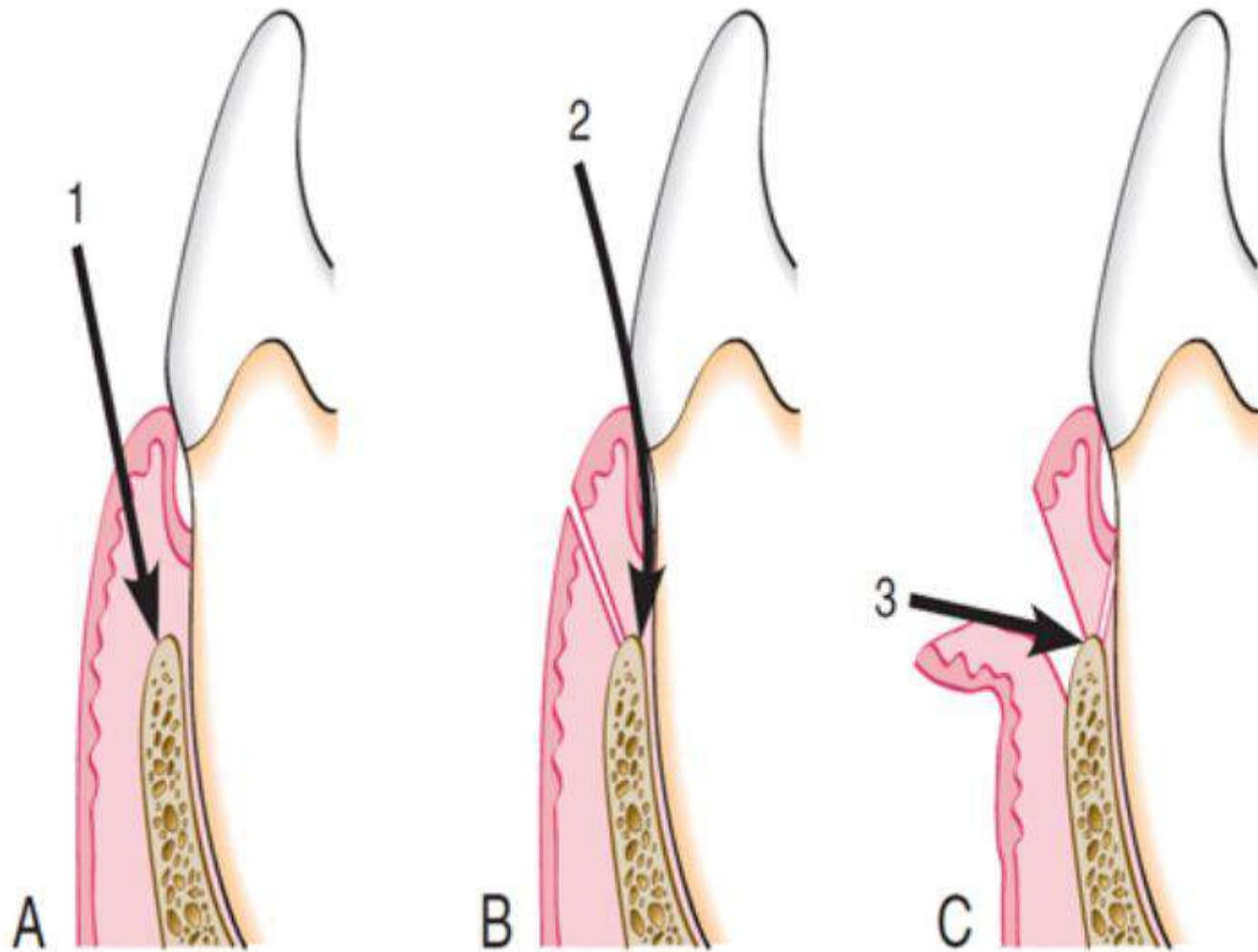
- This incision is carried around each tooth, between the hard tooth structure and the diseased pocket epithelium, to the depth of the junctional epithelium. The 12B scalpel is used.



d:Third incision:Third incision-horizontal

- The horizontal incision is carried along the alveolar crest thus separating the infiltrated tissue from healthy supporting connective tissue, specially in the interdental area. The incision also permits atraumatic removal of the diseased tissue.





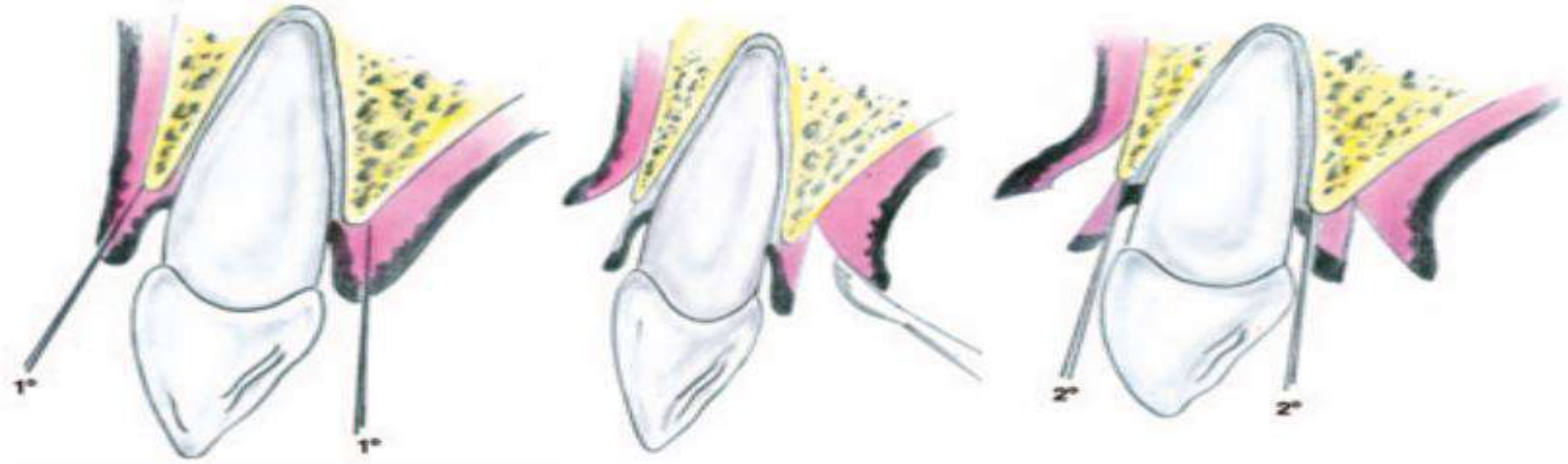
Three incisions of Modified widman flap

e:Direct root planing:Root planing with direct vision

- Fine curettes are used to remove remnants of pocket epithelium and granulation tissue, calculus necrotic cementum to obtain smooth, hard, clean surface.
- Root planing is performed with repeated rinsing.
- Root planing is the most important part of both the modified Widman procedure and all other periodontal surgical procedures.

f: Suturing: Complete coverage of interdental defects

- The labial and palatal flaps are closed over the interdental areas without tension, using interrupted sutures. The flaps should be adapted to the underlying bone and the necks of the teeth. New papillae were created by the scalloped form of the initial incision. These make it possible to cover interdental defects (e.g. bony defects) even when the interdental space is wide. For this reason, placement of a periodontal dressing is not absolutely necessary.





Preoperative clinical view



Internal bevel incision is placed (facial view)



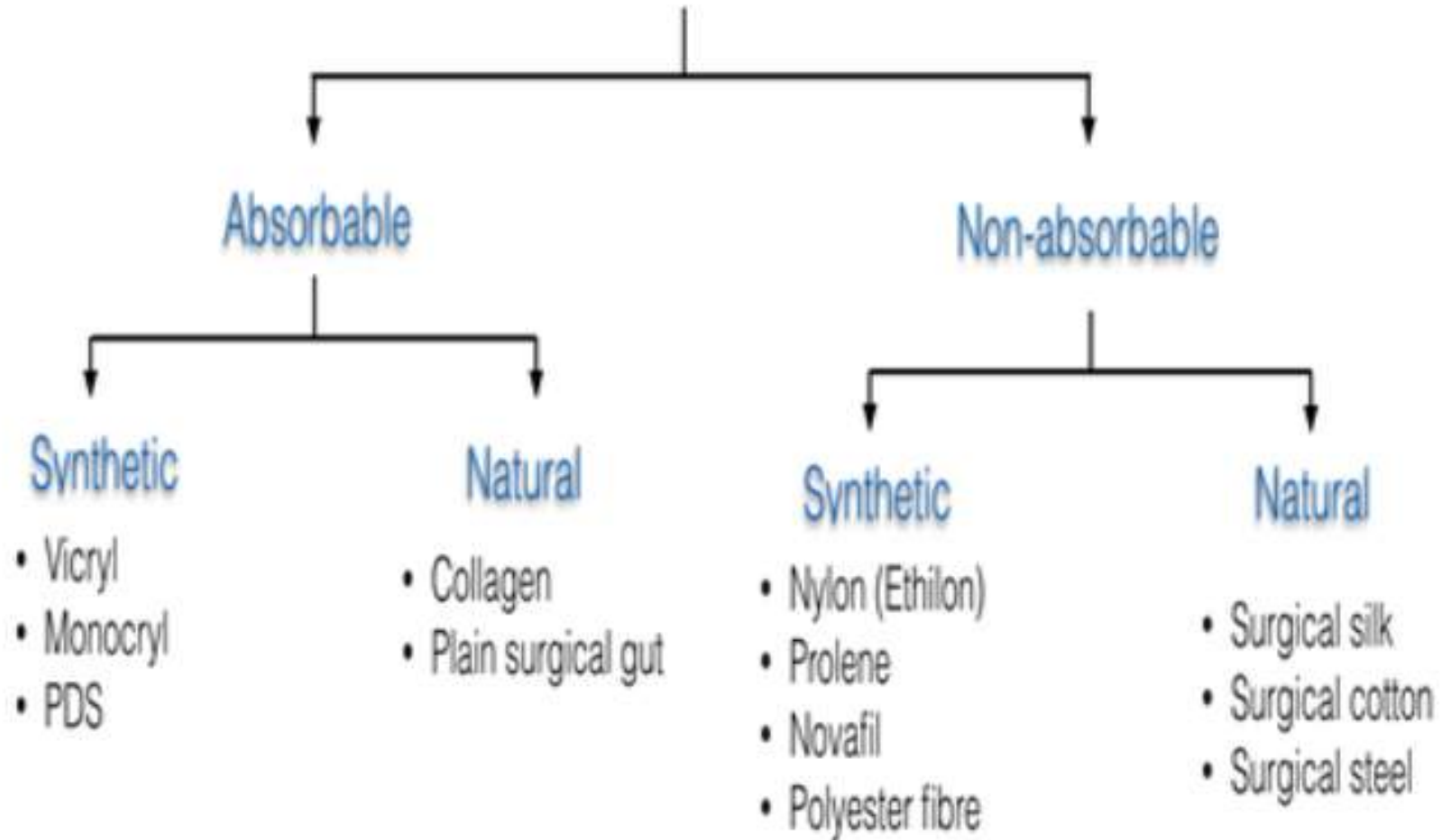
The flap is elevated
Wedge of marginal tissue not yet removed



After thorough debridement



Suture Classification



The Modified Widman flap Advantages

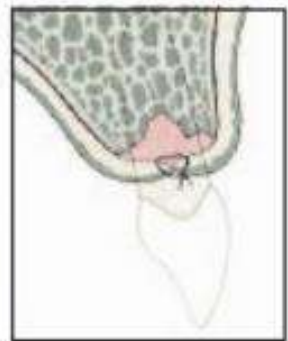
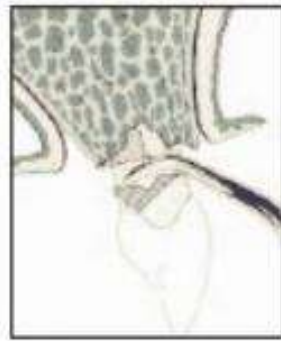
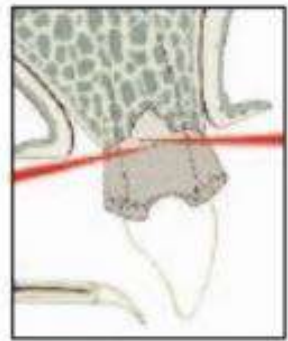
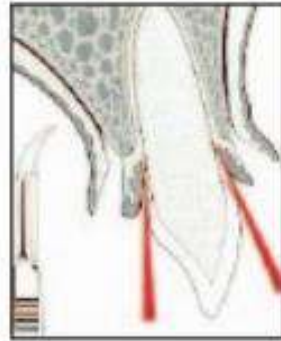
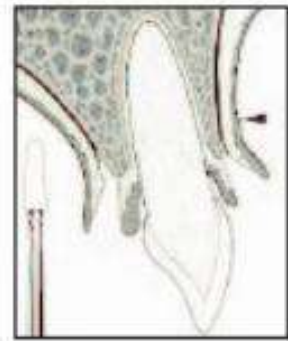
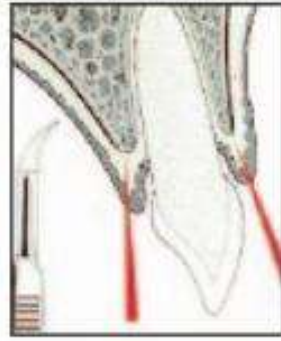
- 1-Good access to root surface to facilitate S+ RP as well as the removal of the pocket epithelium and the inflamed connective tissue.
- 2-Width of keratinized gingiva is maintained.
- 3-Replacement of the flap at presurgical location leads to less exposure of the root surfaces thus minimizes problem of aesthetic (especially anteriorly) and root hypersensitivity.
- 4-Cause minimal amount of trauma to the periodontal tissues and discomfort to the patient.

-
- 5-the possibility of obtaining a close adaptation of the soft tissues to the root surfaces.
 - 6-provides better access to re-establish proper contour of the alveolar bone as well as the potential for bone regeneration in sites with angular bony defect.
 - 7-furcation areas can be exposed.

Healing after flap surgery

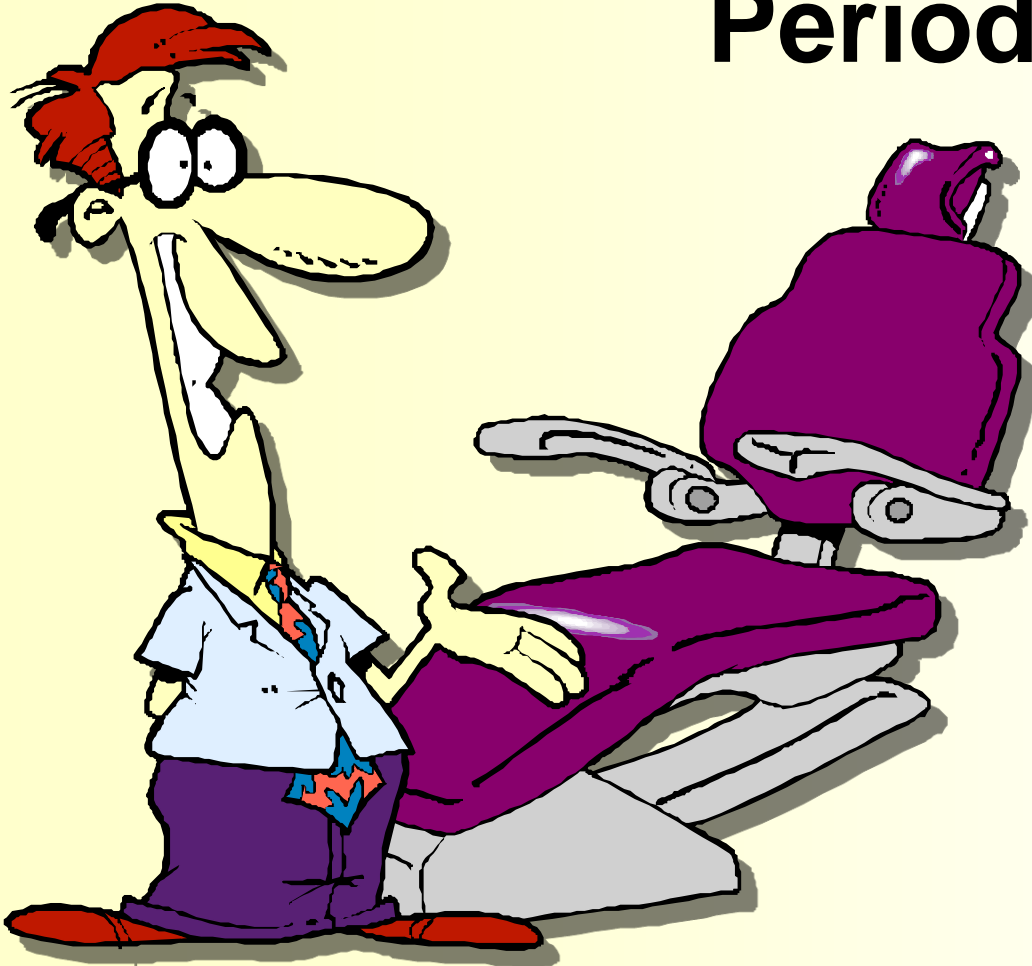
- Following flap procedures and the removal of plaque, calculus and chronically inflamed granulation tissue, healing occurs by the formation of a Long junctional epithelium, this lead to reduced probing depth but that epithelium is more susceptible to plaque induced breakdown than the original connective tissue attachment and consequently post operative plaque control must be a very high standard, a new connective tissue attachment may form following flap procedures, although this cannot be predicted with certainty.

-
- Transient root hypersensitivity and recession of the gingival margins frequently accompany the healing process following close and open S+ RP, thus the patient should be awarned that these results may happen.



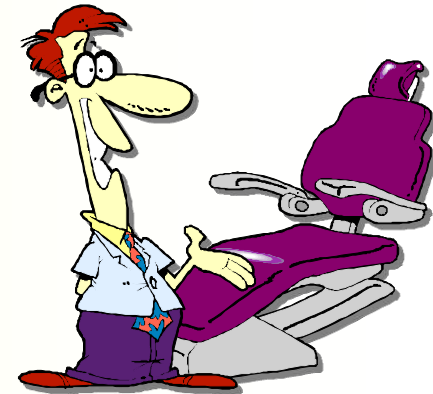
Thank you

Antibiotics in Periodontology



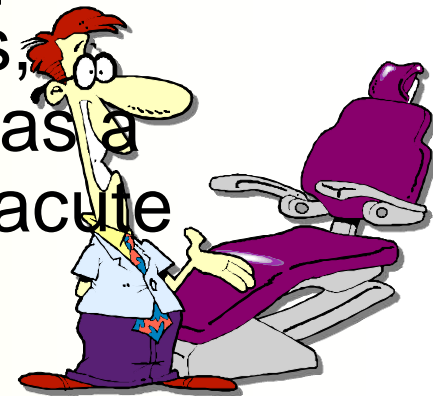
د. سها أسود دهش العزاوي
B.D.S, MSc. Periodontology

- The various periodontal diseases result from susceptible hosts having their periodontal tissues colonized by specific oral pathogens in numbers sufficient to overwhelm their tissue defenses.
- Clinical success in the treatment of these diseases requires reduction of the bacterial load or enhancement of the host tissues' ability to defend or repair itself.



Traditionally, the foundations of clinical success include -

- Education of patients in daily oral hygiene
- Non surgical and surgical mechanical root debridement to remove subgingival bacteria and their accretions from root surfaces
- Supportive periodontal therapy generally at 3-6 month intervals
- Adjunctive Chemotherapy – in certain types of periodontitis like aggressive periodontitis, refractory periodontitis and periodontitis as a manifestation of systemic diseases and acute necrotizing ulcerative gingivitis.

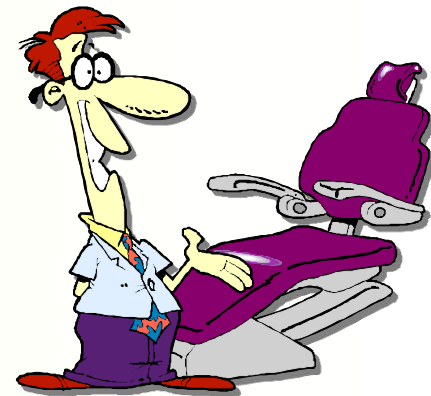


- Antibiotics can be administered
 1. Orally.
 2. Parenterally.
 3. locally.
- In either case, their purpose is to reduce the number of bacteria present in the diseased periodontal pocket.
- Systemic antibiotics may be necessary adjuncts in controlling bacterial infection because bacteria can invade periodontal tissues, making mechanical therapy alone sometimes ineffective.

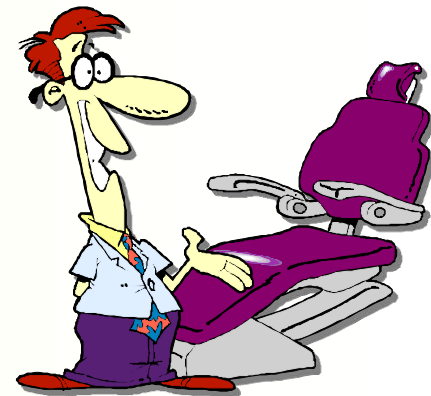




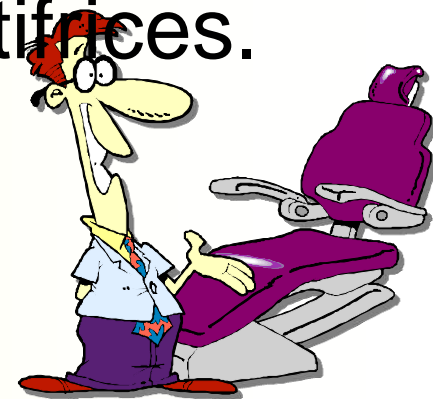
Definitions



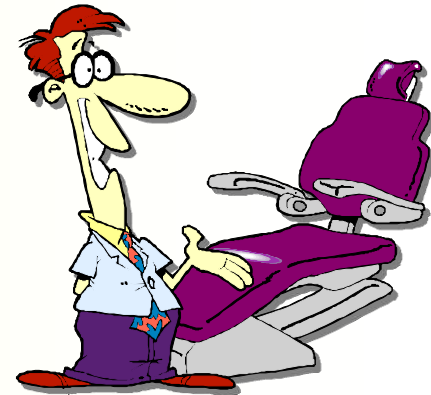
- **ANTIBIOTICS:** are naturally occurring, semi-synthetic or synthetic type of antimicrobial agents that destroy or inhibit the growth of selective microorganisms, generally at low concentrations.



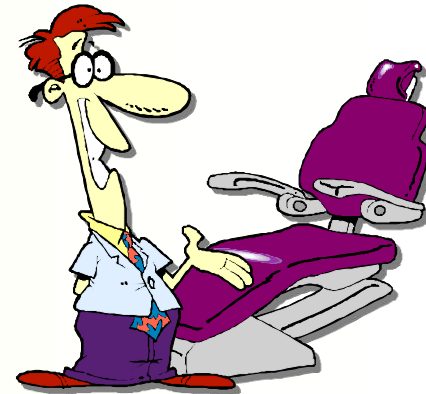
- **ANTISEPTICS:** are chemical antimicrobial agents that are applied topically or subgingivally to mucous membranes, wounds, or intact dermal surfaces to destroy microorganisms and inhibit their reproduction or metabolism.
- In dentistry, antiseptics are widely used as the active ingredients in anti-plaque and anti-gingivitis mouth rinses and dentifrices.



Antibiotics in Periodontal Disease



- ❖ **Penicillins**- Inhibit cell wall synthesis.
- ❖ **Tetracyclines & Macrolides**-Inhibit protein synthesis.
- ❖ **Metronidazole**- Interfere with DNA function.

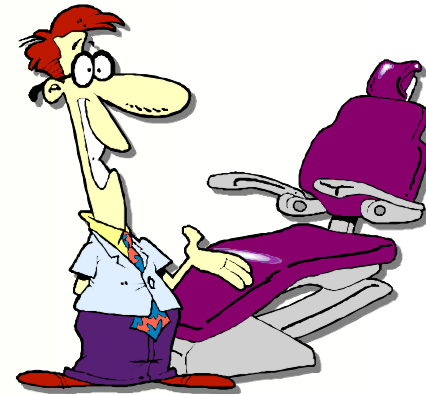


PENICILLINS

- Penicillin was the first antibiotic to be used clinically.
- The present source is *Penicillium chrysogenum*.

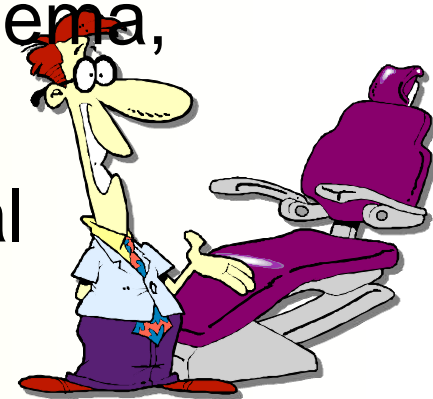
Mechanism of action

- Interfere with cell wall synthesis



Adverse effects

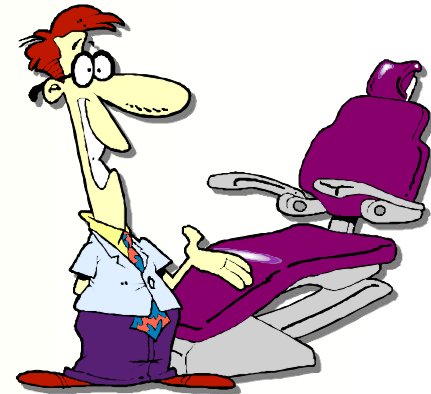
- **Hypersensitivity** – major problem in Penicillin use.
 - Incidence – 1-10%
 - Most common drug implicated in drug allergy
 - Manifestations include fever, rash, itching, urticaria, wheezing, angioneurotic edema,
 - Anaphylaxis is rare but fatal
 - All forms can cause; parenteral > oral



SEMISYNTHETIC PENICILLINS

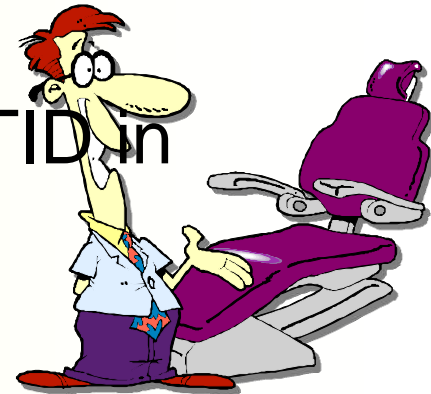
Aim: To overcome shortcomings of Penicillin:

- Hypersensitivity.
- Susceptibility to Penicillinase.
- Poor oral efficacy.
- Narrow spectrum of activity.



Amoxicillin

- Extended spectrum Pn – in addition to gram positive organisms, its also effective against some gram negative organisms.
- Oral absorption is better than Ampicillin; food does not interfere with absorption; higher and more sustained blood levels are produced.
- Incidence of diarrhoea is less.
- Plasma $t_{1/2} = 8$ hr
- Dose: 500mg TID IN ADULTS, 250mg TID in children.



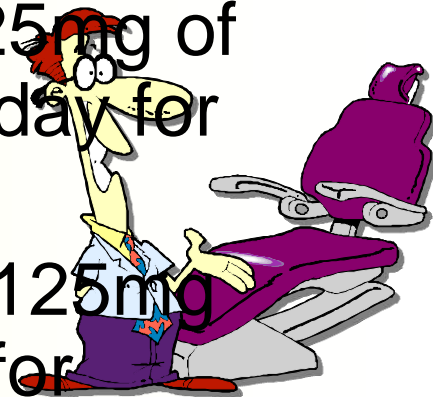
• Augmentin

- It is a combination of amoxicillin with clavulanate potassium.
- Useful in patients with aggressive periodontitis or refractory periodontitis
- Dosage

Augmentin 375mg– amoxicillin 250 mg+125mg of potassium clavulanic acid three times a day for 1 week

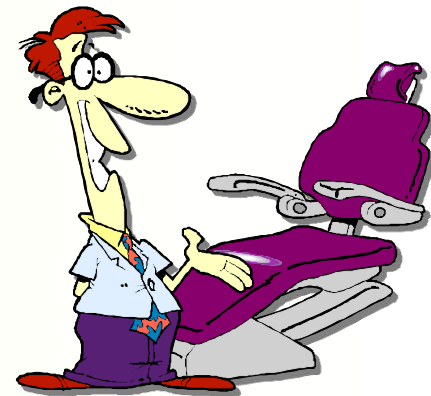
Augmentin 625mg– amoxicillin 500 mg+125mg of potassium clavulanic acid three times a day for 1 week

Augmentin 1000mg – amoxicillin 875 mg+125mg of potassium clavulanic acid twice daily for 1 week

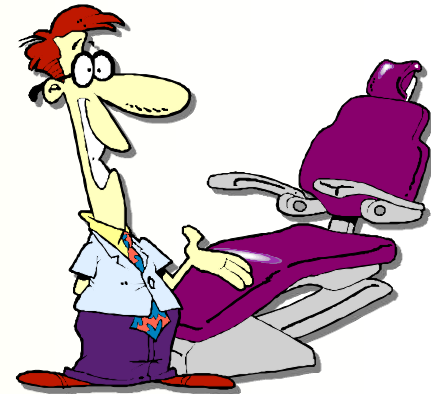


TETRACYCLINES

- Broad spectrum antibiotics
- All are obtained from soil actinomycetes.
- The tetracyclines are primarily bacteriostatic; they inhibit protein synthesis by binding to 30S ribosomes in susceptible organism.



- Widely used in treatment of periodontal diseases
Especially in refractory periodontitis and LAP.
- Concentrate in periodontal tissues and inhibit the growth of microorganisms especially *Aggregatibacter actinomycetemcomitans* (A.ac)
- Concentration of 2 to 10 times in GCF
- Also has anticollagenolytic activity > inhibit tissue destruction.



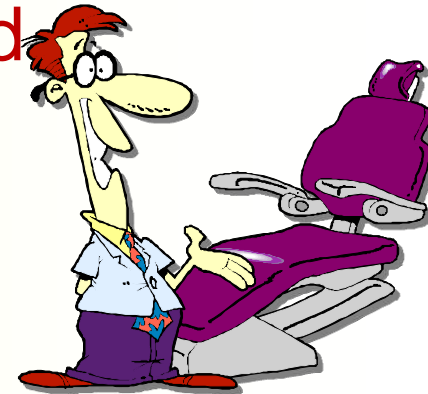
Administration

- Oral capsule is the dosage form in which TCs are most commonly administered.
- 250mg QID
- It should be taken $\frac{1}{2}$ hr before or 2 hrs after food.
- **TETRACYCLINES** have chelating property and form insoluble, unabsorbable complexes with Ca and other minerals. Milk, Iron preparations, non-systemic antacids & sucralfate reduce their absorption.



Effect on bones and teeth

- Tetracyclines have chelating property and Ca-TC chelate gets deposited in developing teeth and bone.
- Brown discoloured ill-formed teeth which are more susceptible to caries
- Temporary suppression of bone growth – deformities and reduction in height.
- **Contraindicated for pregnant women and children.**



Synthetic derivatives:

Doxycycline:

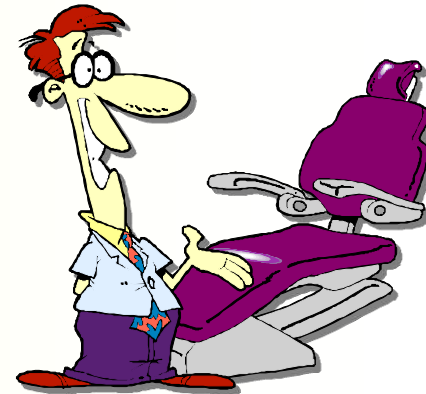
100mg Once daily after loading dose of 200mg

No alteration in absorption unlike Tetracyclines >
no chelating action with calcium

Minocycline:

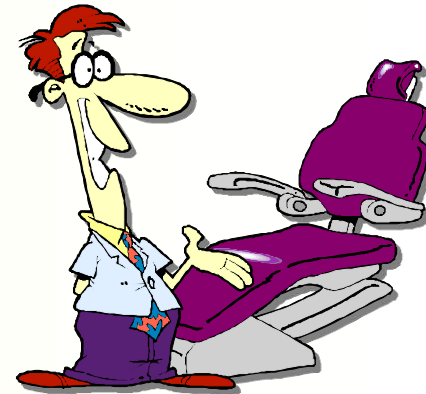
200mg daily

Less photo toxicity and renal toxicity than
tetracyclines

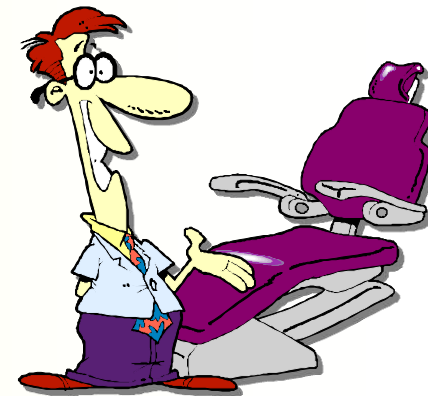


METRONIDAZOLE

- It is primarily an anti-protozoal drug.
- It is selectively toxic to anaerobic microorganisms.
- Inhibits A.ac when given along with amoxicillin.
- Used in treatment of ANUG, LAP and refractory periodontitis.

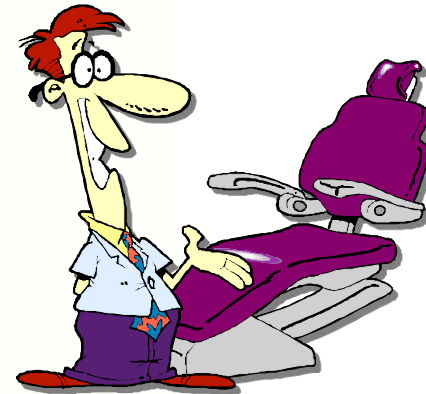


- Plasma $t_{1/2} = 8$ hrs
- Contraindicated in 1st trimester of pregnancy, blood dyscrasias, chronic alcoholism.
- Interaction: adverse reaction to **alcohol** leading to nausea, vomiting, flushing, dizziness, throbbing headache, chest and abdominal discomfort.
- Dose: 250, 500 mg tab TDS



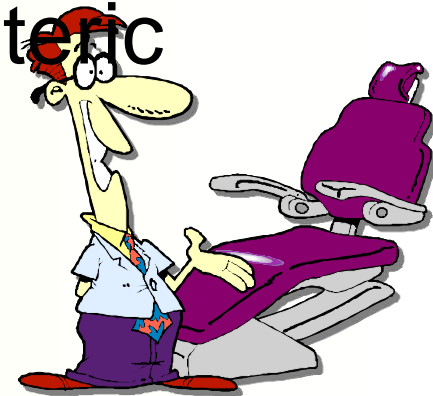
MACROLIDES

- This group includes:
 - Erythromycin
 - Roxithromycin
 - Clarithromycin
 - Azithromycin
 - Spiramycin



Erythromycin

- Bacteristatic at low but cidal at high concentrations.
- It acts by inhibiting bacterial protein synthesis.
- Dose: 250-500 mg QID
- Erythromycin is acid labile and to protect it from gastric acid it is given as enteric coated tablets.



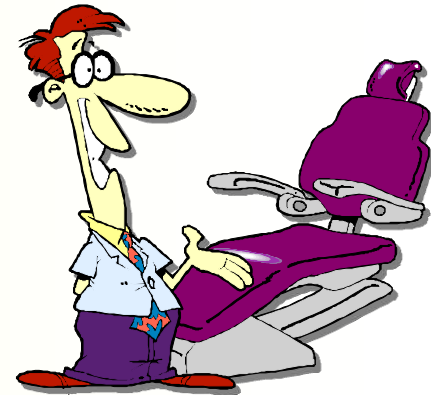
Azithromycin

- Azithromycin has an expanded spectrum,
- It has acid stability, rapid oral absorption, marked tissue distribution and intracellular penetration.
- Concentration in most tissues exceeds that in plasma. Particularly high concentrations are obtained inside macrophages and fibroblasts.
- $T_{1/2} > 50$ hr because of slow release from intracellular sites; Dose: 500 mg OD for 3 days.
- Because of higher efficacy, better gastric tolerance and convenient once a day dosing, Azithromycin is now preferred over Erythromycin.

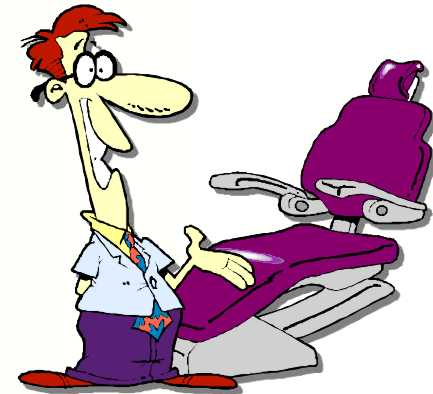




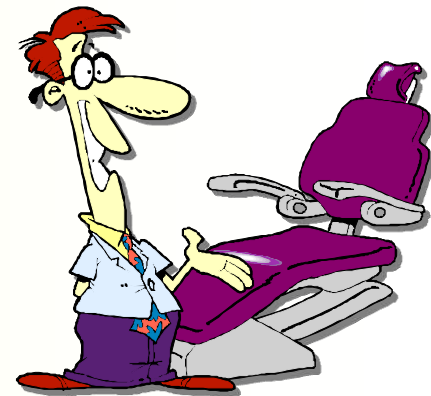
Local Drug Delivery



- ✓ Due to lot of adverse effects with systemic antibiotics– locally delivered agents came into existence.
- ✓ Local administration of drugs directly into the pocket provides greater concentration of drugs directly and reduces the side effects.

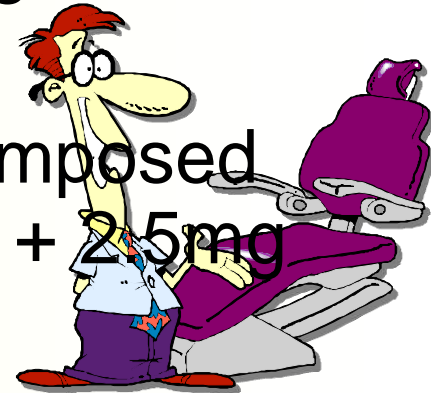


- Local therapy may allow the application of antimicrobial agents at levels that cannot be reached by the systemic route.
- Local therapy may be particularly successful if the presence of target organisms is confined to the clinically visible lesions, whereas systemic administration may reach widely distributed microorganisms.



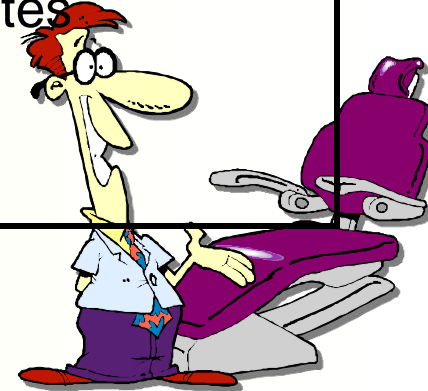
Types of locally delivered agents

- 1- **Tetracycline containing fibers (Actisite)**: copolymer fiber containing tetracycline packed into pocket.
- 2- **Doxycycline (Atridox)**: Gel system available in syringes.
- 3- **Minocycline (Dentamycin & Periocline)**: Gel system available in syringes.
- 4- **Metronidazole (Elyzol)**: Applied in viscous consistency.
- 5- **Chlorhexidine (Periochip)**: Small chip Composed of biodegradable hydrolyated gelatin matrix + 2.5mg chlorhexidine

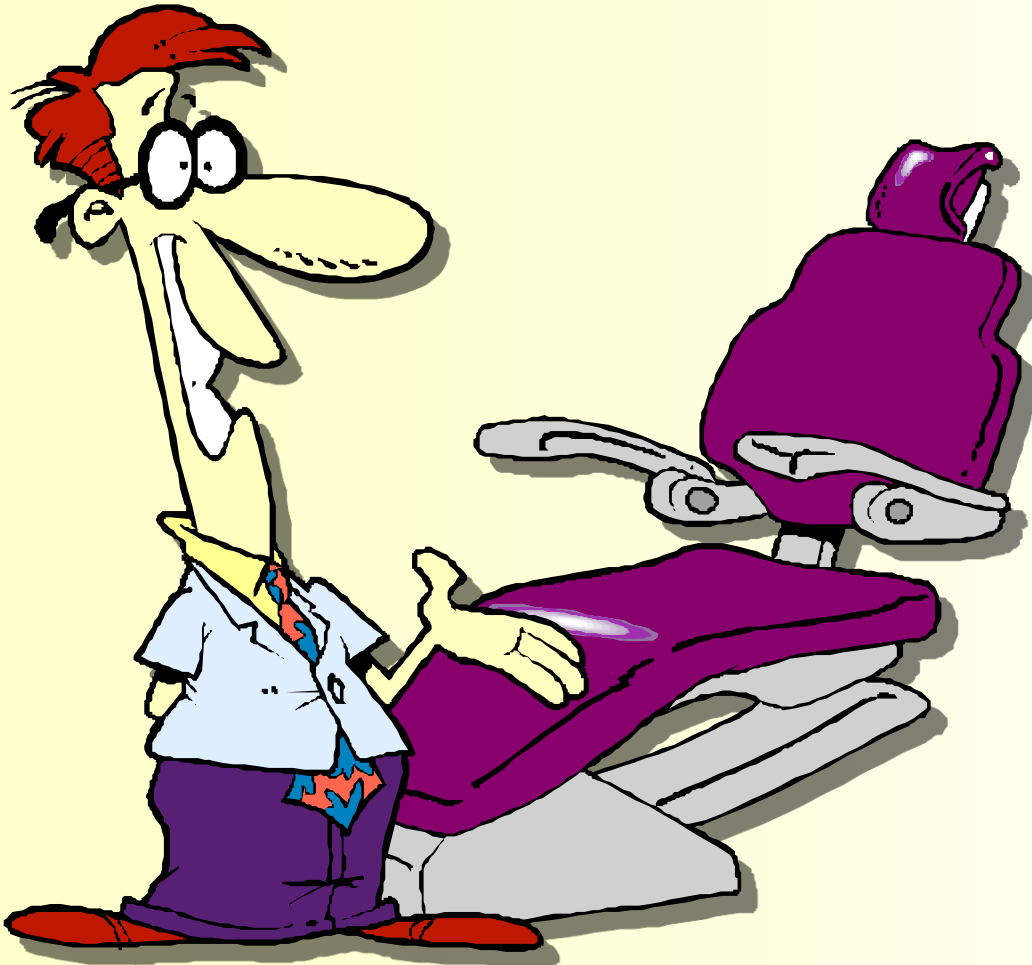


Comparison of local and systemic antimicrobial therapy

	SYSTEMIC	LOCAL
Drug distribution	Reach widely distributed microorganisms	Act locally
Drug concentration	Given at lower concentrations	Applied at higher concentrations at treated sites,
Problems	<ul style="list-style-type: none">-Systemic side effects-Requires good patient compliance	Reinfection from non-treated sites



Thank you



بِسْمِ اللّٰهِ الرَّحْمٰنِ الرَّحِیْمِ

اعداد:

د. نور صباح رحيم

Systemic risk factors

Systemic risk factors can be divided into :-

1-modifiable

- specific microbiota.
- smoking.
- diabetic mellitus.
- stress.
- obesity.
- oral hygiene.
- immunodeficiency.
- drugs (certain medication).
- diet.
- osteoporosis.
- other systemic diseases.

2-non-modifiable

-race.

-age.

-genetic.

-hormonal influence (such as some hormones related to pregnancy).

1-Dental plaque and oral hygiene (microorganisms)

- the primary etiological factor in the development and initiation of periodontal diseases is **dental plaque**.
- dental plaque is a bacterial aggregation on the teeth or other solid structure in the oral cavity.
- dental calculus** which mineralized dental plaque is consider as secondary etiological factor of periodontal disease and it is covered with dental plaque so it serve as reservoir for bacterial plaque (**retentive area**).
- oral hygiene can favorably influence the ecology of the microbial flora in shallow to moderate pocket, but it does not affect host response. Oral hygiene alone has a little effect on sub-gingival micro flora in deep pocket and personal oral hygiene practices among health professional have been shown to be unrelated to periodontitis in these individuals.

Specific microorganisms:-

-although there is sufficient evidence that accumulation and maturation of plaque biofilm is necessary for the initiation and progression of periodontal diseases, studies show that bacterial species colonizing the gingival pocket play variable roles in the pathogenesis of these diseases and may therefore possess different levels of risk of periodontal tissue loss. The subgingival microflora in periodontitis can harbor bacterial species but only a small number has been associated with the progression of diseases such as

-*Aggregatibacter actinomycetemcomitans* (A.a)

-*P. gingivalis*.

-*P. intermedia*.

-*T. forsythia*

These m.o. produce **endotoxins** that will cause severe periodontal destruction.

Tobacco Smoking:

- Tobacco smoking is a major risk factor for increasing the prevalence and severity of periodontal destruction . It was found that the increased risk for periodontitis in smoker was 2.5-7 times greater than non smoker.
- The smoker appear have less gingival inflammation and less bleeding in the gingiva may be explained by decreased gingival vascularity, which includes decreased vascular density, reduce lumen area of gingival vessels (increased vasoconstriction).
- Studies suggested that nicotine increase rate of proliferation of gingival epithelium which can contribute to the reduction of inflammatory clinical signs in the gingival tissues. These are physiological effect of smoking on the etiology of periodontal disease , with decreased gingival crevicular fluid flow.
- Smoking suggest an imbalance between bacterial challenge and host response which may due to changes in the composition of subgingival plaque with increase in the number and or virulence of pathogenic organisms.

The microbiological effect of smoking: •

1- it is increase the colonization of shallow periodontal pockets by periodontal pathogens and increase levels of periodontal pathogens in deep periodontal pocket. •

2-smokers may have higher level of *Tannerella forsythia*, *P.gingivalis* and *T. denticola*. •

3- smoke derived aryl hydrocarbons and bacterial LPS may act additively to inhibit bone formation , may explain why periodontal bone loss is greater and bone healing is less successful in smokers than non smokers with periodontal infections. •

The immunological effect of smoking: •

1-Nicotine causes decrease immune response and impair PMNs chemotaxis and phagocytosis. •

2-Increase the production of TNF-alpha , IL6 , these immune mediators are known to lead to more sever destructive inflammation in the periodontal tissue. •

3-Reduction in the serum concentration of Immunoglobulin as IgGs which is essential in the production against periodontal infection. Also smoking decrease the level of salivary IgA antibodies. •

----The risk of periodontal disease increase with number of cigarettes smoked per day. smoking of cigars and pipes carries the same risk as smoking cigarettes. Exposure to secondhand smoke may be associated with an increased risk for developing periodontal disease. •

Diabetes mellitus:

- Diabetes is a modifiable factor in the sense that though it cannot be cured, it can be controlled.
- studies have been done which suggest that poorly controlled diabetes respond less successfully to periodontal therapy relative to well-controlled and non-diabetic.
- it is a complex metabolic disease characterized by chronic hyperglycemia.
- uncontrolled diabetic is associated with a reduction in the defense mechanisms (neutrophil dysfunction, impairment of chemotaxis and phagocytosis), atherosclerosis and reduce normal gingival flow, increased susceptibility to infection include periodontitis and poor wound healing..
- Diabetic mellitus does not cause gingivitis or periodontitis , but it alters the response of periodontal tissues to local factor.
- Diabetic patients with poor oral hygiene may have very sever gingival inflammation , deep periodontal pockets, rapid bone loss, and frequent periodontal abscess.

Neutropenia:•

-The diagnosis of Neutropenia is based on clinical signs and • symptoms as well as absolute neutrophil count a relatively deficiency in neutrophil number can dramatically increase susceptibility to infectious disease.

-Neutropenia is considered clinically significant when the neutrophil • count falls below 1,000 cell/ml (normal adult range: 1,800-8,000 cell/ml).

-Disorders of neutrophil function also increase the host susceptibility • to infection, by decrease chemotaxis , migration, phagocytosis and killing.

Osteoporosis: •

-many of the studies conducted to date suggest there is a relationship • between skeletal osteoporosis and bone loss to the extent that postmenopausal osteoporosis may result in dental osteopenia (reduction in bone mass due to imbalance between bone formation and resorption) involving the jaw , and particularly the mandible. Osteoporosis was significantly associated with severe alveolar bone loss and the prevalence of periodontitis causes in post menopause women and in men with advancing age.



Drugs (medications):

-Gingival enlargement is a well –known consequence of the administration of some drugs as anticonvulsants (Phenytoin or Delantin), immunosuppressant (Cyclosporine) and Ca channel blockers (Antihypertensive drugs) such as

-Amelodipine.

-Diltiazam .

-Nicardipine.

-Nifedipine.

Nisoldipine. -

-Verapamil -



Obesity:

-Obesity is one of the most significant health risks of modern society . A condition associated with obesity (the metabolic syndrome) , this a clustering of dyslipidemia and insulin resistance may exacerbate periodontitis. Obesity has been postulated to reduce blood flow to the periodontal tissues, promoting the development of periodontal diseases. Furthermore, obesity may enhance immunological or inflammatory disorders , which might be the reason obese subjects tend to exhibit escalating poor periodontal status relative to non-obese individuals.

Psychological factors:

- Studies have been demonstrated that individuals under psychological stress are more likely to develop clinical attachment loss and loss of alveolar bone.
- One possible link in this regard may be increased **glucocorticoid** secretion that can depress immune function , increase insulin resistance , increase in production of **IL-6** in response to increased psychological stress and potentially increased risk of periodontitis. Another study suggest that host response to *P. gingivalis* infection may be compromised in psychological stress individuals also the relationship is simply due to the fact that individuals under stress are less likely to perform regular good oral hygiene and prophylaxis.

Nutrition:

-The potential mediating role of nutrition in the oral health-systemic diseases relationship has increased interest in the effect of nutrition in oral health and periodontal disease. However, efforts to correlate the nutritional deficiency to periodontal disease have yielded conflicting results. Vitamins are coenzyme required for metabolism and health.

HIV/AIDS

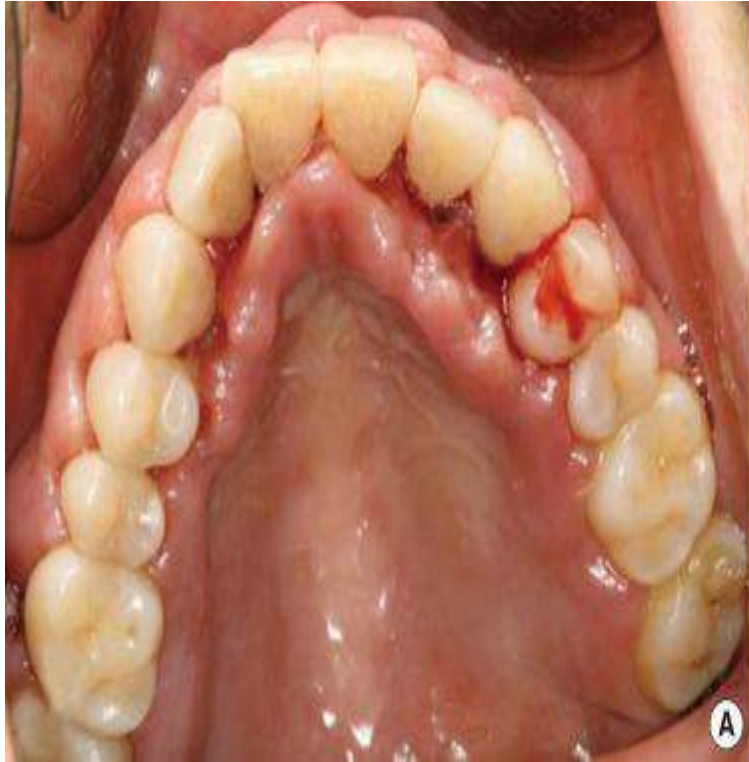
It has been stated that the immune dysfunction (immunosuppression) associated with human immunodeficiency virus (HIV) infection and acquired immunodeficiency syndrome (AIDS) increases susceptibility to periodontal disease. Those patients often had severe periodontal destruction characterized by necrotizing ulcerative periodontitis.



Hematological factors •

Hemorrhagic gingival overgrowth with or without necrosis is a • common early manifestation of acute leukemia. Patients with chronic leukemia may experience similar but less severe periodontal changes. Chemotherapy or therapy associated with bone marrow transplantation may also adversely affect the gingival health.







Non –modifiable systemic risk factors: •

- Genetic •
- Female Hormonal Alteration •
- Race •
- Gender •

Genetic factors

There is evidence that genetic differences between individuals may explain why some patients develop periodontal disease and other do not. Genetic factor may play an important role in determining the nature of the host response and may effect the function of phagocytic immune cells or the structure of the epithelium or connective tissue.

-One of the disease is **papillon- Lefevre syndrome** which is a rare hereditary disease characterized by hyperkeratotic skin lesion in the palms, soles, knee and elbow and severe destruction to the periodontium with early loss of primary and permanent teeth.

-Other disease is **aggressive periodontitis** which has familial aggregation (which may be seen in one family) . Some immunological defects are associated with aggressive periodontitis. The periodontal disease has a link with some genetic factor by:

1-A specific interleukin (**IL-1**) genotype has been associated with severe chronic periodontitis.

2-Neutrophil abnormalities are under genetic control.

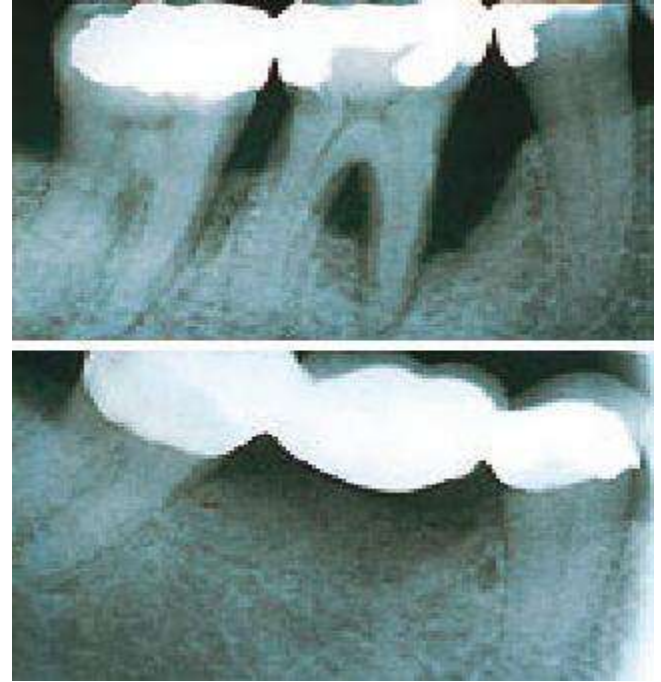
3-Genetic play a role in regulating the titer of protective **IgG2** antibody response to **A.a** in patients with aggressive periodontitis.



الطبيب | سلسلة العيادات

12 - متلازمة فرط التقرون الراضي الأحمصي
Papillon Lefevre syndrome





Pregnancy, puberty, menopause(hormonal) (Female • hormonal alteration):

- -Pregnancy associated gingivitis is inflammation of the gingival tissues associated with pregnancy. This condition is accompanied by increase in steroid hormones in GCF and increase in the levels of (P. intermedia) bacteria which use steroid as a growth factors. The increase in sex hormones may exaggerate the inflammatory response to dental plaque which means small amount of plaque may lead to gingivitis.
- -Puberty is also accompanied by an exaggerated response of the gingiva to local irritation.
- -During menopause, estrogen deficiency will reduce bone mineral density.



GINGIVA IN PREGNANCY

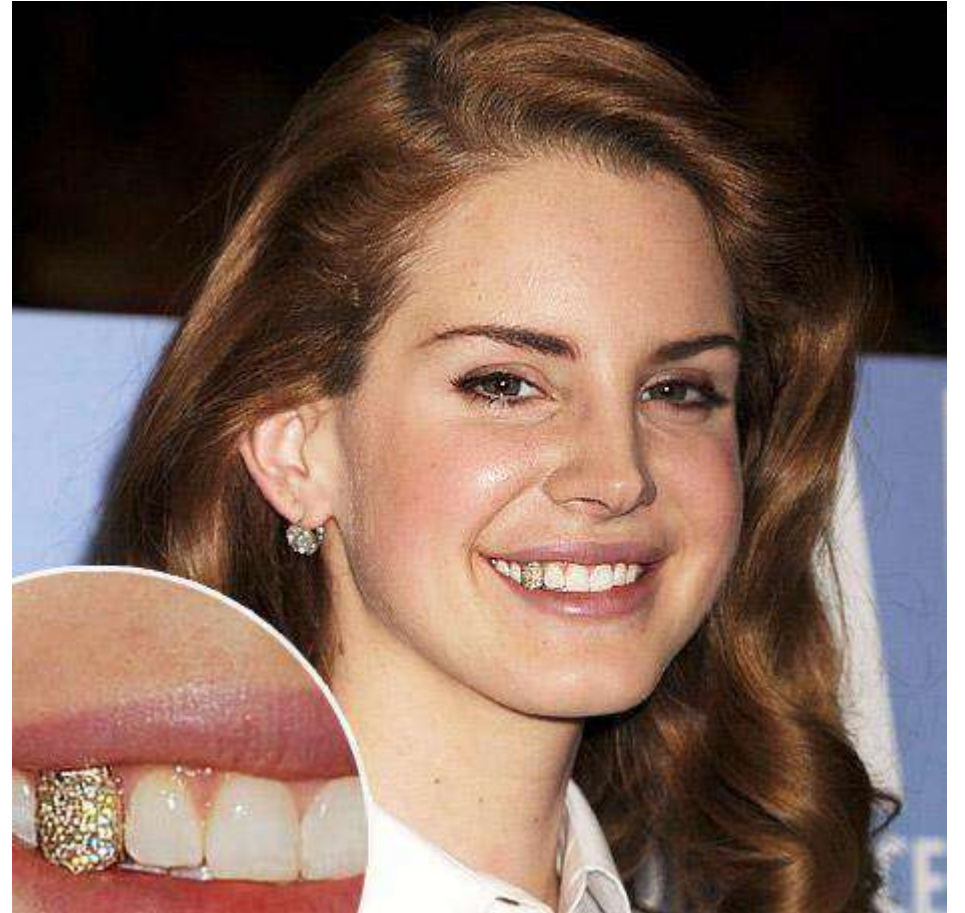


Gender •

-Numerous studies reported higher periodontal destruction among **males** • **compared to the female** population. The reason for these sex differences are not clear, but they are thought to be related to **ignorance of oral hygiene**, which is usually observed among **males**.

Race: •

-the level of attachment loss is also influenced by **race/ethnicity**, although the • exact role of this factor is not fully understood . Certain racial/ethnic groups, particularly subjects of **African and Latin American background** , have a **higher** risk of developing periodontal tissue loss than other group.





THANK YOU.

TREATMENT PLANNING OF PERIODONTAL DISEASES

PRESENTED BY:

DR: NOOR SABAH IRHAYYIM

Aim of the treatment planning:

-To coordination of all treatment procedures for the purpose of the establishment and maintenance of a well-functioning dentition in a healthy periodontal environment such as:.

1-reduction or resolution of gingivitis(bleeding on probing BOP)

2-reduction of probing pocket depth (PPD)

3-elimination of open furcation in multi-rooted teeth.

4-absence of pain

5-individually satisfactory esthetic and function

- Bacterial specificity and pathogenicity as well as the •
disposition of the individuals for diseases (local and systemic
resistance) may influence :

1-onset •

2-rate of progression •

3-clinical characteristic of plaque associated dental disorder •

because both caries and periodontal diseases represent
opportunistic infections associated with biofilm formation on
the surface of teeth.

Treatment of patients divided into 4 phases:

1-systemic phase of therapy. •

2-intial phase (cause –related therapy) •

3-correction phase (additional therapeutic measures) •

4-maintenance phase(supportive periodontal therapy) •

SYSTEMIC PHASE OF THERAPY:

- 1-precaution for the protecting the general health of the dental team and other patients against infections and contagious disease (such as hepatitis, HIV, Herpes simplex virus and TB), the minimal precaution (wearing rubber gloves, masks, protective glasses, the dental team should be vaccinated against hepatitis)
- 2-protective of the patients health against harmful systemic effects of routine therapy, the complications most commonly encountered in the dental office are (infection, bleeding, cardiovascular incidence, allergic reaction, control anxiety and low pain threshold, smoking, treatment of emergency).

1-infection: patients with cardiac disorder involving the endocardium susceptible to infection with infective endocarditis by blood born infection by do extraction, scaling, root planning, periodontal surgery, implant leading to bleeding and bacteremia. So that prophylaxis antibiotic 1 hr. before dental procedure.

2-bleeding: patients with anticoagulant drug (Salicylate), liver cirrhosis, high alcohol consumption, hemophilia have high risk to develop bleeding, so consultation with patients physician and treatment in small segment.

3-Cardiovascular disease:

A-patient with anticoagulant drug such as aspirin have bleeding tendency

B-patients with cardiovascular drugs(antihypertensive. Diuretic, anti arrhythmic) may increase hypotensive episodes, or develop angina pain or congestive heart failure as result of stress, so control anxiety and pain and do short procedures.

4-allergic reaction and drug interaction: most common allergic to local anesthesia (Novacain), penicillin, sulfa derivative, so ask to replace this drug.

5-controlling anxiety and low pain threshold:

A-valium in night before, in the morning, and half an hour before extensive treatment

B-apply local anesthesia.

C-post operative analgesic such as voltaren, ponstan.

Objectives of initial phase (cause-related therapy):

- 1-motivation the patients to combat dental disease (patient information) •
- 2-giving the patients instruction on the proper oral hygiene techniques (self-performed plaque control methods) •
- 3-scaling and root planning. •
- 4-antimicrobial therapy (local or systemic) •
- 5-control or elimination of additional retention factors for plaque such as : •
correction of restoration and prosthetic irrational factors and excavation of caries and restoration.
- 6-occlusal therapy •
- 7-orthodontic treatment •

MOTIVATION:

Detailed information must be given to the patients regarding • his/her periodontal diseases , its etiological factors symptoms , consequence, prognosis and the relationship between the presence of dental plaque and calculus in the mouth and the location of sites showing dental diseases by using plaque disclosing agents.

Disclosing agents:

Is a chemical compound (tablet or solution) that stains • dental plaque such as erythrosine, fuschcin or fluorescein,



Brushing:

The most widespread mean of actively removing plaque at home is tooth brushing. The efficacy of brushing with regard to plaque removal is dependent on:

1-the design of brush •

2-the skill of individuals using of brush(manual dexterity) •

3-frequency and duration of brushing •

4-the morphology of the dentition(crowding, spacing, gingival phenotype, longer teeth, exposed dentin, ...)

Oral hygiene instruction component:

1-self assessment •

2-self examination •

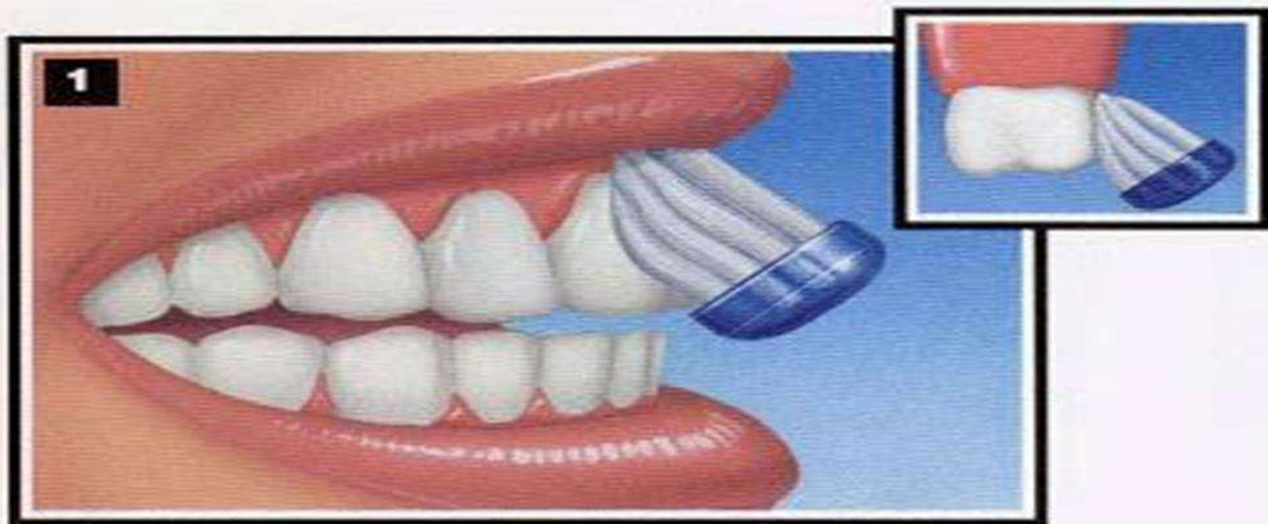
3-self monitoring •

4-self instruction •

For this purpose disclosing agent is applied before brushing, •
and the aid of mirror, the patients can identify the amount of
plaque formed after last brushing and give information
about his/her cleaning performance.

Methods of tooth brushing

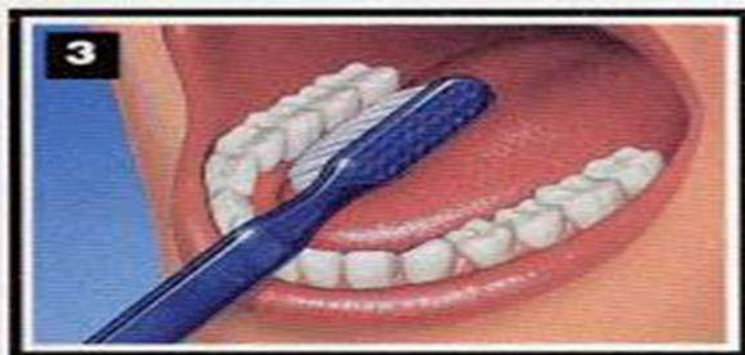
- 1-horizontal brushing (scrub) •
- 2-vertical brushing (leonard technique) •
- 3-circular brushing (fones technique) •
- 4-vibrotary technique(stillman technique) •
- 5-roll technique (modified stillman) •
- 6-charter technique •
- 7-sulcular technique (Bass technique) •
- 8-modified bass technique •



Place bristles along the gumline at a 45° angle. Bristles should contact both the tooth surface and the gumline.



Gently brush the outer tooth surfaces of 2-3 teeth using a vibrating back, forth & rolling motion. Move brush to the next group of 2-3 teeth and repeat.



Maintain a 45° angle with bristles contacting the tooth surface and gumline. Gently brush using back, forth & rolling motion along all of the inner tooth surfaces.



Tilt brush vertically behind the front teeth. Make several up & down strokes using the front half of the brush.



Place the brush against the biting surface of the teeth & use a gentle back & forth scrubbing motion. Brush the tongue from back to front to remove odor-producing bacteria.



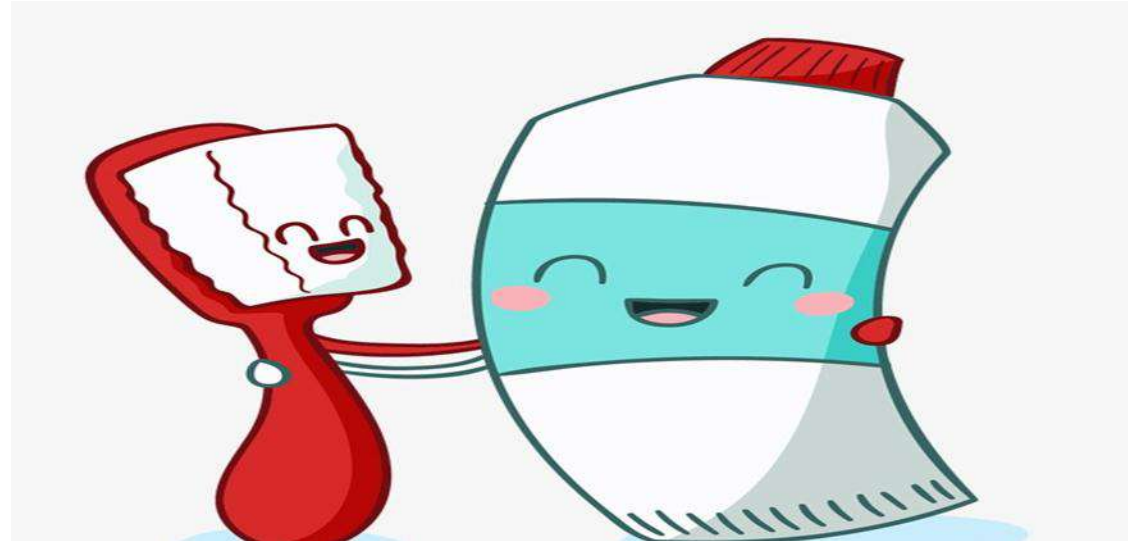
Tooth brushing requirement:

- the features of manual toothbrush in periodontics must be **Nylon, • Soft-medium, rounded ends filaments, the small head** is easier to reach all areas of oral cavity and should be trimmed flat and must be multi-tufted with same length.
- the **3 brush head** clean vestibular, occlusal, oral tooth surfaces this • design is superior to other types of brush.



Frequency of brushing: 2 in a day , just before going to bed •

Duration: for a minimum 2 minutes, tooth brush should be replaced every 2-3 months, because a worn toothbrush with frayed filaments loses resilience and is less effective in removing plaque •



Electric toothbrushes:

Used mainly: •

1- elderly •

2-those with arthritis in their hands and wrists •

3-children •

4-hospitalized individuals •

5-physically or mentally handicapped •

6- neurological disorders •

7-consider easier and faster than manual. •



Dentifrices:

-used in combination with tooth brush for plaque removal, should have: •

1-fluoride: prevent caries •

2-desensitizing agent.: for exposed dentin •

3-anti-plaque agents(Triclosan, Stannous fluoride, chlorhexidine) •

4-anti-calculus agent •

5-bicarbonate , reduce the acidity of dental plaque •

6-cleaning and polishing agent (abrasive agent) •

7-whitening agents •

8-detergents •



gettyimages®
kickstand

185406309

Dental Floss:

- have a variety of thickness and type •
- have with or without wax •
- large interdental space use a thicker type than thinner one •
- 40 cm floss used wind around middle finger, 10cm between middle •
finger, 3cm between thumbs
- using a sawing movement •
- avoid fast movement to avoid damage of gingiva •
- do not worry if gum is bleed slightly after first one use , this stop •
after number time of use,

- **unwaxed** used in normal tooth contact •
- **waxed** used in tight proximal tooth contact •
- floss holder to facilitate flossing to use •
- **Tap**: type of broad dental floss used to clean bridge pontics •
- **Super floss**: used for crown, bridge and orthodontic appliance •
- floss used in vertical direction not horizontal to avoid development of grooved surface •
- flossing is difficult and time consuming •



Woodsticks:

- made from wood and have a triangular cross section and a vailable in •
different thickness
- hold between thumbs and first finger •
- don't worry if gum bleed from the first cleaning with wood stick , this •
disappear after a repeat using of it.



Single tufted tooth brush

- used to clean can not reach by other type of brush •
such as locks, wire brace of orthodontic, groove
- back surface of last molar, long standing tooth. •
- used also in fixed appliance (crown and bridge) •

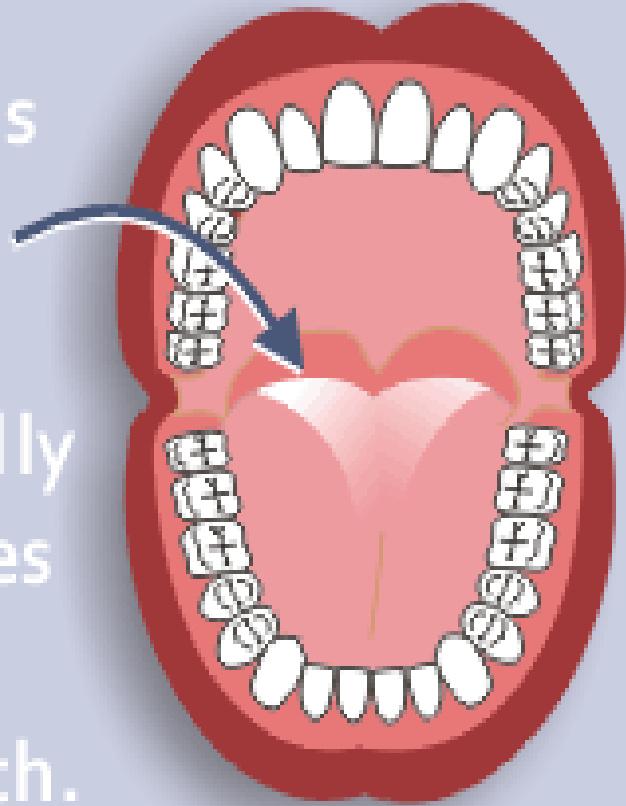




Tongue cleaner:

- called **scraper** •
- gag reflex develop after using •
- remove bacteria between tongue groove •
- most effective one is loop shaped •
- clean mostly middle and first part •
- used mainly in patients with halitosis •

It's the accumulation
of
debris
here
that
usually
causes
bad
breath.



© 2013 WMD S, Inc.

Animated-Teeth.com

Animated-Teeth.com

Bacteria and
Debris



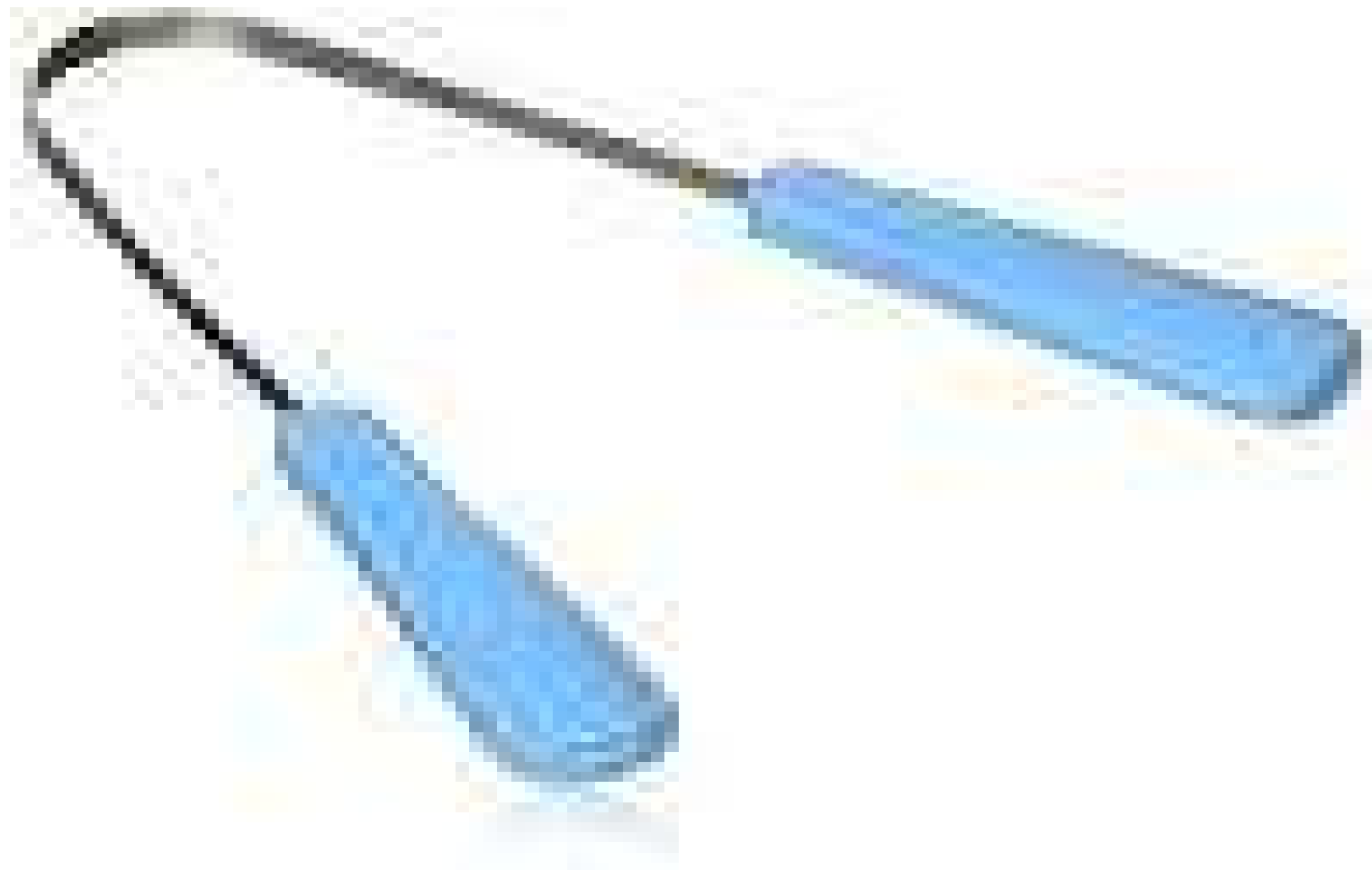
© 2013 WMD S, Inc.



Animated-Teeth.com



© 2013 WMD S, Inc.



Inter dental brushes

Used when we have: •

- widely open interdental spaces •
- when root surface with concavities or groove have been exposed •
- in through –and-through furcation defects in periodontal patients •
- have different size and shape, most common forms are cylindrical, conical shaped head •
- can be used for carry fluoride or chlorhexidine gel •
- if is not properly used can be lead to dentin hypersensitivity •
- used without dentifrices •



Dental water jet

- The daily use of oral irrigation has been shown to reduce gingivitis and bleeding
- -the pulsating hydrodynamic forces produced by irrigator can rinse away food debris from interdental spaces and plaque retentive area
- -use adjunct to brushing and flossing, and used with water or chlorhexidine to improve plaque inhibition and had anti-inflammatory effect

An effective Interdental Cleaner



THANKYOU •