Papillon - Lefevre Syndrome - A Brief Review of Diagnosis & Management

Dr T. Dinesh Kumar¹, Dr Ajay Kumar Pillai², Dr Parimala Kulkarni³, Dr Swapnil Moghe⁴, Dr Vineesh Vishnu⁵, Dr Saurabh Dhanraj Yadav⁶

¹Reader, Dept. of Maxillofacial Pathology, R.V.S Dental College & Hospital, Coimbatore, India
², ⁴Reader, Dept. of Maxillofacial Surgery, Peoples Dental Academy, Peoples University, Bhopal, India.
³H.O.D. Dept of Peadodontics & Preventive Dentistry, Peoples Dental Academy, Peoples University, Bhopal, India.
⁵Post graduate student, Dept. of Public health Dentistry, Peoples Dental Academy, Peoples University, Bhopal, India.
⁶Post graduate student, Dept. of Peadodontics & Preventive Dentistry, Peoples Dental Academy, Peoples University, Bhopal, India.

I. Introduction:

Papillon lefevre syndrome is a rare autosomal recessive disorder with a characteristic clinical features of palmoplantar hyperkeratosis with severe early onset periodontal destruction followed by the premature shedding of both the deciduous and permanent teeth⁴,⁵. It was first described by two French physicians papillon MM and Lefevre in 1924.

II. Etiology

The cause of Papillon lefevre syndrome is not well understood¹⁰,¹⁵. Recent studies indicate a cathepsin C gene mutation localized to chromosome 11q 14-21 is responsible for Papillon lefevre syndrome¹³,²³. The cathepsin c-gene encodes a cysteine lysosomal protease also known as dipeptidyl amino peptidase I or cathepsin C. Cathepsin C is capable of removing dipeptides from the terminus of protein substrates but at higher pH it also exhibits dipeptidyl transferase activity. The cathepsin C gene is expressed in epithelial regions commonly affected by Papillon lefevre syndrome such as palms, soles, knees and keratinized oral gingiva¹⁶,²¹. The pathological clinical findings associated with Papillon lefevre syndrome suggest that cathepsin C is functionally important in structural growth and development of skin in susceptibility to periodontal disease⁸. It is unknown if the profound periodontal disease susceptibility is a consequence of altered integrity of junctional epithelium is eliminated, the severe gingival inflammation resolves⁴⁰. A variety of immunological findings have been reported in Papillon lefevre syndrome patients, including decreased monocyte chemotaxis, decreased neutrophil chemotaxis, impaired neutrophil phagocytosis, altered superoxide production and decreased response, but it has been difficult to extrapolate results of these studies¹².

III. Clinical manifestations:

The Papillon lefevre syndrome has a reported incidence of 1-4 cases per million person and both the sexes are equally affected and there is no racial predominance⁷. The age of onset is generally between 1-4 years⁴,⁶,²¹.

The cardinal signs of Papillon lefevre syndrome are, palmar plantar hyperkeratosis and early onset periodontitis. In affected subjects, the development and eruption of the primary teeth proceed normally, but the eruption of these into the oral cavity is associated with gingival inflammation and subsequent rapid destruction of the periodontium. After exfoliation, the inflammation subsides and the gingiva resumes a healthy appearance, although the third molars are sometimes spares⁸,⁹,¹⁰.

It can be associated with hyperhidrosis of palms and soles, resulting in a foul smelling odour, psoriasis form, scaly plaques on the elbows and knees, ectopic calcifications in choroid plexus and tentorium, follicular hyperkeratosis, hearing loss, recurrent infections, mental retardation and retinitis pigmentosa⁴,⁵,⁹,¹⁴,²¹. We report a case of 15 year old male patient presented with the chief complaint of multiple loss of teeth. Clinical history revealed that he had normal emergence of deciduous teeth at 8-9 months of age, which started loosening at three years and were all eventually lost by four to five years of age. Patient or his guardians were not sure about the time of eruption of permanent teeth, but described gingival bleeding during brushing and eating, after the eruption of permanent teeth. There was loosening of permanent teeth from 12 years of age and eventually two permanent teeth were lost by 15 years of age. Bleeding was also associated at the time of tooth
loss. With the complaint of bleeding patient visited a local dentist who performed oral prophylaxis. Patient was normal for a couple of months but again noticed bleeding gums and then visited to our department.

The patient was moderately built with a steady gait. His physical and mental development was also normal. Medical history was noncontributory. Parents were not of consanguineous marriage. In pregnancy history of mother, she had full term uneventful pregnancy. Dermatological examination revealed dry skin with normal development of hair and nails. There were symmetric, well-demarcated, brownish, keratotic, plaques affecting the skin of his palms on the dorsal surface and soles on the ventral surface as seen in (Figure 1). Nail changes such as transverse grooving and fissuring are apparent in advanced cases.

![Figure 1: Keratotic Plaque on the dorsal surface of both the hand.](image1)

Intra Oral examination shows premature exfoliation of permanent maxillary and mandibular teeth. Generalized periodontitis was noticed. Prompt bleeding on probing was also noticed.

![Figure 2: Intra Oral examination shows premature exfoliation of permanent maxillary teeth.](image2)

![Figure 2: Intra Oral examination shows premature exfoliation of mandibular teeth.](image3)
Panoramic view showed generalized loss of alveolar bone, complete loss of bone support around all the present teeth.

IV. Investigation and diagnosis:
- Radiological feature is severe destruction of alveolar bone which gives the teeth a “floating in air” appearance and could be associated with intra-cranial calcifications.
- Histopathological findings have consisted of hyperkeratosis, occasional patches of parakeratosis, acanthenosis and a slight perivascular inflammatory infiltrate.
- Differential diagnosis includes Hiam-munk syndrome. Both Papillon lefevre syndrome and Hiam-munk syndrome are allelic variants of cathepsin C gene mutation. Both are classified as type IV palmo-planter-ectodermal keratoderma. The clinical distinction of Hiam-munk syndrome from Papillon lefevre syndrome being atropic changes of nails and arachnodactyly. Some suggest, Hiam-munk syndrome is a clinical variant of Papillon lefevre syndrome. Some times cutaneous lesions of Papillon lefevre syndrome, may be mistaken for psoriasis unless they are correlated with oral dental findings.

V. Treatment:
A multidisciplinary approach is important for the case of patients with Papillon lefevre syndrome. It is usually not necessary to treat cutaneous lesions unless they interfere with patient activities. The skin manifestations are usually treated with emollients salicylic acid and urea may be added to enhance either effects. Periodontal treatment may consist of frequent scaling and oral hygiene instructions. Oral retinoids including acitretin, etertinate and isotretinoin are the main stay to the treatment of both the keratoderma and periodontitis associated with Papillon lefevre syndrome. Treatment may be more beneficial if it is started during the eruption and maintained during the development of the permanent teeth. Antibiotic therapy when incorporated delays the shedding of the teeth. Early extraction of teeth too has been advocated to prevent bony loss. Moreover this allows solid base for subsequent use of artificial dentures.

A course of antibiotics should be tried to control the active periodontitis in an effort to preserve the teeth and to prevent bacteremia and subsequent pyogenic liver abscess. The risk of pyogenic liver abscess should be kept in mind in evaluating these patients when they present with fever of unknown origin.

VI. Summary:
Papillon lefevre syndrome is a rare, autosomal recessive disorder occurring between the first and fifth years of life, which is characterized by the palmo-planter keratoderma with periodontitis followed by the premature shedding of both deciduous and permanent teeth. The cause is not well understood but recent studies have reported that loss of function mutations affecting both the alleles of the cathepsin C gene. Further research at molecular level carries significant implications for understanding the susceptibility of the disease.

Take Home Message
1. History of normal eruption of the primary and permanent Teeth.
2. Inflamed and swollen gingivae.
3. Bleeding on probing on slight provocation.
4. Mobility of teeth, with deep infra bony pockets.
5. Severe alveolar bone loss.
7. Lesions subside with exfoliation of teeth.
8. Hyperkeratotic papules present on the hands and soles.

References
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