LITERATURE REVIEW

Oral and periodontal features of autosomal recessive syndromes: a tabular review

Características oral e periodontal de síndromes autossômicas recessivas: uma revisão tabular

Sesha REDDY¹, Kharidhi Laxman VANDANA², Shishir Ram SHETTY³
1 – Department of Periodontics – College of Dentistry – Gulf medical university – Ajman – United Arab Emirates.
2 – Department of Periodontics – College of Dental Sciences – Davangere – India.

doi: 10.14295/bds.2019.v22i2.1713

ABSTRACT

Objective: To systematically review the data and results of case reports of autosomal recessive syndromes associated with periodontitis. Material and Methods: An internet search using Google and PubMed search engine and keywords- autosomal recessive, periodontitis, syndromes, periodontium and gingiva was carried out. Full-text articles in the English language of all the case reports and reviews that were published in journals from the year 1966 to 2016 were obtained and evaluated and presented in tabular form. Abstracts and articles published in other languages were not included in the review. Results: The data available from the clinical trials were analyzed and presented under broad headings of, systemic features, dental features, periodontal features and laboratory findings presented in tabular form. Conclusion: Many autosomal recessive syndromes with dental component also present with changes or alteration in the periodontium thus stressing the fact that thorough periodontal examination is important during the medical evaluation of patients with syndromes.

KEYWORDS
Syndromes; Autosomal recessive; Periodontitis; Periodontium; Gingiva.

RESUMO

Objetivo: Analisar sistematicamente os dados e resultados de relatos de caso de síndromes autossômicas recessivas associadas à periodontite. Material e Métodos: realizou-se uma pesquisa na internet usando os sites Google e PubMed com as palavras-chave: autossômica recessiva, periodontite, síndromes, periodonto e gengiva. Os critérios de inclusão foram restritos aos artigos em texto completo em língua inglesa, relatos de casos e revisões publicados em periódicos de 1966 a 2016. Resultados: Os dados extraídos de cada estudo foram agrupados da seguinte forma: as síndromes associadas a características sistêmicas, aos achados dentários e aos achados periodontais, apresentados no formato de tabelas. Conclusões: Diversas síndromes autossômicas recessivas que apresentam alterações dentárias também podem apresentar alterações no periodonto, ressaltando assim, o fato de que o exame periodontal completo é importante durante a avaliação médica de pacientes com síndromes.

PALAVRAS-CHAVE
Síndromes; Autossômica recessiva; Periodontite; Periodonto; Gengiva.
INTRODUCTION

A syndrome is defined as the collection of signs and symptoms associated with any morbid process and constituting together the picture of the disease. [1] Multiple manifestations could be caused by underlying developmental or metabolic conditions. Diagnosing and distinguishing a syndrome from other pathologies is challenging for a clinician. All syndromes have genetic components. Mutations in various classes of genes lead to craniofacial, dental and oro-facial defects. [1-5].

Some are more obviously important than others depending on the age of onset of the disease, degree of mental or physical impairment, number of affected persons and cost of care. Although many of these disorders are not preventable or curable early detection may allow significant improvement in health care. [2-5] Dentist is usually in a position to recognize a previously unrecognized genetic or birth defect problem in a patient or family. Since many syndromes affect the oral structures in a unique way it aids in diagnosis. Although treatment of genetic diseases is not yet available, maintenance of good oral health is important. [6]

Syndromes are classified as autosomal dominant, autosomal, recessive or sex-linked.

Identification of these syndromes and genetic counseling plays an important role in the prevention of progression of these syndromes to the next generation. In ideal situations of an autosomal dominant case, it is possible to trace the disorder through successive generations of the family and on an average equal number of males and females will be affected. In some conditions, the disease may skip generations in these situations the normal individual is assumed to have inherited the mutant gene but do not express it. This is referred to as incomplete penetrance. [1] In other cases, there may be a number of abnormalities with different individuals within a family exhibiting one or more of these abnormalities. This is referred to variable expression of the trait and it is possible to have a condition which exhibits both incomplete as well as variable expression. The periodontium is affected in some of the syndromes and identification of periodontal manifestations plays a vital role in the diagnosis of syndromes itself. In this review, an attempt has been made to review autosomal recessive syndromes of periodontal interest. In autosomal recessive inheritance, two copies of a disease allele are required for an individual to be susceptible to expressing the phenotype. (With an autosomal recessive condition, a gene alteration needs to be present in both copies of a particular gene to cause sufficient impairment to cell function to cause disease. The alterations are located on an autosome. A person with an autosomal recessive condition must have inherited one gene alteration from each parent. In autosomal recessive inheritance, people with one copy of the gene alteration do not have the condition. They are said to be carriers for the autosomal recessive condition.[2-6] Typically, the parents of an affected individual are not affected but are gene carriers.

Many of these cases are rare and thus the evidence comprising is mainly on case reports rather than epidemiological studies. However, it is relevant to consider syndromes background to these severe periodontal cases as it may eventually be relevant to treat and also to understand the pathogenesis of periodontitis in these patients.

METHODOLOGY

An internet search using Google and PubMed search engine and keywords- autosomal recessive, periodontitis, syndromes, periodontium and gingiva was carried out. Inclusion criteria were restricted to full-text articles in English language, case reports and reviews that were published in journals from the year 1966 to 2016. The articles were reviewed and presented in tabular form. Articles published in languages other than English were not included in this review. Abstracts were not included in the review. The data extracted from each study were grouped as follows: syndromes associated with systemic features, dental and periodontal findings are presented in the tabular format. [Table 1]
Table 1 - Autosomal recessive syndromes with systemic, dental, and periodontal features

<table>
<thead>
<tr>
<th>Sl</th>
<th>Syndrome</th>
<th>Systemic Features</th>
<th>Dental Features</th>
<th>Periodontal Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Alport Syndrome (AS) [7]</td>
<td>Hematuric nephropathy, renal failure, hearing loss, ocular abnormalities and changes in the glomerular basement membrane of the lamina densa.</td>
<td>No dental features reported</td>
<td>Gingival Hyperplasia</td>
</tr>
<tr>
<td>3</td>
<td>Cross Syndrome (CS) [2-6]</td>
<td>White hair, blond skin; melanocytes decreased with reduced tyrosine activity; mental retardation; very rare.</td>
<td>No dental features reported</td>
<td>Gingival and alveolar enlargement,</td>
</tr>
<tr>
<td>4</td>
<td>Chediak–Higashi syndrome (CHS) [9-11]</td>
<td>Partial oculocutaneous albinism usually involving the skin, eyes and hair. Susceptibility to infection, Photosophobia, frequent pyogenic infections and lymphadenopathy.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Göhlich-Ratmann Syndrome (GRS) [12]</td>
<td>Mental retardation and epilepsy, brachymetacarpalia, hirsutism, bulbous soft nose, thick floppy ears with abnormal configuration. Brachymetacarpia, Tetralogy of Fallot and the other with congenital hypothyroidism and bilateral ureteral stenosis.</td>
<td>No dental features reported</td>
<td>Gingival hypertrophy</td>
</tr>
<tr>
<td>6</td>
<td>Haim–Munk Syndrome (HMS) [13-15]</td>
<td>Inactivation of cathepsin C, Palmoplantar hyperkeratosis, acro-osteolysis, atrophic changes of the nails, and a radiographic deformity of the fingers. Arachnodactyly (spider fingers), and pes planus (flat foot).</td>
<td>No dental findings</td>
<td>Severe early-onset periodontitis that affects both primary and permanent dentitions, with severe alveolar bone destruction</td>
</tr>
<tr>
<td>8</td>
<td>Kindler Syndrome (KS) [18-20]</td>
<td>Skin fragility, patchy hyperpigmentation, hyperkeratosis of palms and soles, diffuse skin wrinkling, sun sensitivity, eczematoid dermatitis, skin fragility, patchy,</td>
<td>Atrophy of buccal mucosa, limited mouth opening, malocclusion, erosion of hard palate, geographic tongue,</td>
<td>Bleeding, gingival swelling, desquamative gingivitis. Patients present with severe periodontitis of both the primary and secondary dentition, resulting in severe alveolar bone loss and premature exfoliation of the teeth.</td>
</tr>
<tr>
<td>9</td>
<td>Kostmann Syndrome(KS) [21-25]</td>
<td>Congenital neutropenia recurrent bacterial infections early in life severe ear, skin, respiratory and oral infections,</td>
<td>Severe oral infections, persistent gingivitis</td>
<td>Gingival swelling and bleeding, severe inflammation of all gingival tissues and periapical abscesses with mobility, aggressive periodontitis with alveolar bone loss.</td>
</tr>
<tr>
<td>10</td>
<td>Lazy Leukocyte Syndrome (LLS) [26,27]</td>
<td>Severe neutropenia, Recurring infection, deficiency in neutrophil chemotaxis and systemic neutropenia,</td>
<td>Painful stomatitis, gingivitis and recurrent ulcerations of the buccal mucosa and tongue</td>
<td>Periodontitis progressing to the point of advanced alveolar bone loss and tooth loss has been reported. Individuals are susceptible to aggressive periodontitis.</td>
</tr>
<tr>
<td>11</td>
<td>Maroteaux-Lamy Syndrome (muco- polysaccharidosis VI) [MLS] [28,29]</td>
<td>A large head, short neck, corneal opacity, open mouth associated with an enlarged tongue, enlargement of the skull, and a long anteroposterior dimension bone dysplasia, joint restriction, organomegaly, heart disease, and corneal clouding, among several other problems, and reduced life span.</td>
<td>Unerupted dentition, dentigerous cyst follicles, malocclusions, condylar defects. High palate, open bite impacted and/or included teeth, thickening of the periocular follicle, and changes in the temporomandibular joint.</td>
<td>Gingival hyperplasia.</td>
</tr>
</tbody>
</table>
RESULTS

The autosomal recessive syndromes discussed within the scope of this review are- Alport syndrome, Bardet-Biedl syndrome, Cross syndrome, Chediak–Higashi syndrome Gölich-Ratmann syndrome, Haim–Munk syndrome, Hurler’s Syndrome, Kindler syndrome Kostmann syndrome, Lazy Leukocyte Syndrome, Maroteaux-Lamy syndrome, Murray-Puretic Drescher syndrome, Papillon–Lefèvre syndrome, Rutherford syndrome and Winchester syndrome. All the above mentioned autosomal recessive conditions have significant dental and periodontal findings as stated in the table. Some of the aggressive periodontal features might lead to tooth loss which in turn may further deteriorate the quality of life in these individuals. On the evaluation of the tabular data, the oral features of the syndromes followed three patterns. In the first pattern, only periodontal findings are observed with no distinct dental features or mucosal features. The syndromes with only periodontal/gingival findings without dental hard tissue findings include Alport syndrome, Cross syndrome, Golich Ratman syndrome, Haim-Munk syndrome, Murray-Puretic Drescher syndrome, Papillon–Lefèvre syndrome, Rutherford syndrome and Winchester syndrome. In the second pattern of clinical features, the syndromes included both hard tissue and periodontal findings. The syndromes in with this feature include Maroteaux-Lamy syndrome, Hurlers syndrome and Bardet-Biedl syndrome. The third pattern observed in the oral features of the syndromes mentioned in the table is a combination of oral mucosal features and periodontal/gingival findings. The syndrome that occurs in this category is Lazy Leukocyte syndrome, Kostmann syndrome, Kindler syndrome and Chediak Higashi syndrome.
DISCUSSION
Apart from classifying the syndromes based on the combination of periodontal/gingival, dental and mucosal findings the syndromes can also be classified under the following categories, namely. The syndromes listed above have been discussed under the three categories namely-

• Syndromes associated with gingival findings- Alport Syndrome, Bardet-Biedl Syndrome, Göhlich-Ratmann syndrome, Maroteaux-Lamy syndrome, Murray-Puretic Drescher syndrome, Ramon Syndrome, Rutherford syndrome.

• Syndromes associated with gingival and periodontal findings- Cross syndrome, Haim–Munk syndrome, Kostman syndrome, Papillon–Lefevre syndrome.

• Syndromes associated with dental, gingival and periodontal findings- Chediak–Higashi syndrome, Hurler's Syndrome, Kindler syndrome, Lazy Leukocyte Syndrome.

Alport Syndrome: The mutation presented in AS produces defects in chains α3, α4, and α5(IV) of Type IV collagen which is an important component of the periodontium. This defect leads to disruptions of the epithelial bonds leading to gingival hyperplasia which appears as a prominent intraoral finding in this syndrome. [38]

Bardet-Biedl Syndrome (BBS): The syndrome is characterized by the six main general clinical features. Oral findings that have been frequently associated with this syndrome include hypodontia, small roots, high arched palate, microdontia and gingival enlargement. [39]

Maroteaux-Lamy syndrome (MLS): The intraoral finding associated with of this syndrome include gingival hyperplasia, hypertrophy of the maxillary alveolar ridge, macroglossia, unerupted dentition, malocclusions and dentigerous cyst-like follicles. [40]

Murray Puretic Drescher Syndrome-JHF: Juvenile Hyaline Fibromatosis is characterized by disproportionate accumulation for hyalin in various tissues such as skin, stomach-intestinal system, heart muscle, surneals, skeletal muscles, spleen, and lymph nodes thyroid tissue. Gingival hypertrophy is very prevalent and distinctive oral finding in this syndrome. Some cases of advanced gingival hypertrophy have caused difficulties in tracheal intubation. Presence of cervical spine and temporomandibular joint contractures further adds to the difficulty in the tracheal intubation procedure. [41]

Ramon’s Syndrome (RS): Gingival fibromatosis (GF) associated with Ramon’s syndrome could be attributed to the adverse effect of the anti-epileptic drug often used as a treatment modality for seizures associated with this syndrome. However, there are cases with sporadic occurrence of fibromatosis often in the absence of seizures or use of antiepileptic drugs. The characteristic feature of occurrence of fibromatosis is its coincidence with the eruption of teeth. [2-5,42,43]

Rutherford syndrome (RS): The association of gingival fibromatosis with corneal opacities and retarded tooth eruption is recognized as an autosomal dominant trait known as Rutherfurd syndrome. [2-5]

Winchester Syndrome (WS): Winchester Syndrome is often attributed to alteration in a gene called MMP2. It is suggested that this condition is caused by a nonlysosomal connective-tissue alteration. The protein inactivation mutation is found on the matrix metalloproteinase 2 gene (MMP2) MM2 which is responsible for bone remodeling. This alteration in the pattern of bone remodeling leads to periodontal tissue destruction. [37,44]

Cross Syndrome (CS): Cross syndrome is, almost certainly, an autosomal recessive disorder characterized by gingival fibromatosis, microphthalmia, mental retardation, athetosis, and hypopigmentation. [2-5]
Haim-Munk Syndrome (HMS): Haim-Munk syndrome is an exceedingly rare autosomal recessive disorder characterized clinically by palmoplantar hyperkeratosis, aggressive periodontitis with severe alveolar bone destruction. The other key features of the syndrome are onychogryphosis, pes planus, arachnodynctly, and acro-osteolysis. A definite link to the history of consanguinity seems to be responsible in all documented cases. Presence of typical dermatological, periodontal, and radiological features are generally important for the diagnosis of this syndrome. A classic distinguishing feature between HMS and Papillon–Lefever syndrome [PLS] is that the severity of the periodontal destruction is less severe in later. [14,42] Early diagnosis seconded by prompt intervention often leads to better retention of permanent teeth.

Papillon–Lefever syndrome: The key features of PLS consist of severe gingivostomatitis and periodontitis. No changes have been reported in the literature regarding eruption pattern or sequence associated with primary teeth. However morphological alteration such as microdontia, root resorption, and incomplete root formation have been reported frequently. Vertical bone loss at a younger age is the common radiographic feature observed in most of published case reports. [46]

Chediak–Higashi syndrome: Severe gingivitis and gingival bleeding accompanied by early tooth loss are the commonly reported oral features of CHS. Premature tooth loss in this syndrome could be attributed to alveolar bone loss secondary to inflammation associated with periodontal pathogens such as Prophyromonas gingivalis, Prevotella intermedia and Tannerella forsythia. Aphthous, pyoderma and oral ulcer have sometimes been reported in cases of CHS. [47]

Hurler’s syndrome: Hurler’s syndrome also referred to as mucopolysaccharidosis type IH is an autosomal-recessive inherited disorder. It often signifies the classical prototype of a mucopolysaccharide disorder. [48,49] The oral and dental findings of MPS I-II comprise of hyperplastic gingiva, macroglossia, high-arched palate, short mandibular rami with abnormal condyles, spaced hypoplastic peg-shaped teeth with a retarded eruption. Dental radiographic features such as localized cyst-like radiolucencies often mimicking dentigerous cyst have been reported. [49]

Kindler syndrome: The syndrome is frequently associated with aggressive periodontitis leading to tooth loss. desquamative gingivitis along with gingival bleeding also have been noticed in some case reports. [50]

Lazy leukocyte syndrome: Oral mucosal findings such as excruciating pain associated stomatitis often recurrent ulcerations have been observed. Gingivitis and periodontitis progress to the point of advanced alveolar bone loss and tooth loss has been reported. [51]

**CONCLUSION**

Genetics will endure and dictate dental, medical diagnostics in the future. Human and microbial genomics proteomics and metabolomics added with pharmacogenomics will shape the future of the medical profession. All dentist and physicians need to comprehend the concepts of genetic variability, its interface with the environment and its repercussions for the patient and population healthcare. In the near future genomics will augment preventive, diagnostics and therapeutics. Few reviews articles, in general, have discussed the dental and periodontal features of syndromes. However, in this review, an attempt has been made to discuss very specifically the periodontal and dental features of autosomal recessive syndromes. A number of autosomal recessive syndromes and the details of dental and periodontal features have been discussed in the above review making it a fairly useful article for a clinician or researcher looking out for comprehensive data on this specific topic. The results obtained from the review also aid
in categorizing the syndromes based on the presence of dental periodontal and mucosal features. This categorization reduces the number of syndromes in the list of differential diagnosis and aids in the diagnosis of these syndromes which are rare in occurrence.

REFERENCES


Oral and periodontal features of autosomal recessive syndromes: a tabular review


