SYSTEMIC DISORDERS AFFECTING DENTAL PATHOLOGY

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Abstract: A retrospective overview of systemic disorders which might be associated with dental pathology is made. They are grouped as follows: (a) congenital dental developmental disorders, (b) chromosomal anomalies, (c) radiations, (d) immune disorders, (e) intoxications, (f) neurological alterations, (g) gastrointestinal diseases, (h) osteodystrophy and associated conditions, (i) skin diseases, (j) metabolic and endocrine disorders, (k) craniofacial malformation syndromes and other congenital general malformations. The associated dental pathology is described in each case.

Key words: Dental diseases, Symptoms, Systemic disease, Retrospective overview.

INTRODUCTION

Dental pathology may be divided into five etiological groups /1/ infections (e.g., caries) /2/, traumatisms /3/, disorders resulting from dental wear (e.g., attrition, erosion, and abrasion) /4/, pathologic formations /5/ and dental developmental disorders — the latter usually being associated to other extra oral clinical manifestations that may or may not form part of a common syndrome.

Many diseases and pathological conditions, involving practically all human apparatuses and systems, exhibit associated dental pathology or manifestations. The present study reviews those systemic disorders that may associate dental pathology, grouped as shown below. The grouping of such systemic diseases into categories is complicated; however, since group overlapping inevitably occurs. On reviewing the etiologies of dental disorders, no uniform classification criteria are to be found among the different authors who have addressed the subject. As an example, in the case of morphology — structural dental enamel defects, Neville et al. (1) propose an exhaustive classification comprising trauma to developing tissues, the ingestion of chemical substances, chromosomal anomalies, infections, hereditary diseases, malnutrition, metabolic alterations and neurological disorders. In contrast, Wright (2) describes only three etiological groups: metabolic diseases, syndromic hereditary disorders and nonsyndromic hereditary disorders (e.g., amelogenesis imperfecta and other enamel developmental disorders). As commented above, the classification or grouping of the different diseases poses difficulty — particularly when referring to congenital disorders. In this context, it is accepted that in addition to the etiological factors associated to the dental anomalies described below, other underlying factors — probably genetic, related to tooth development and individual resistance to disease — are also implicated (3).

CONGENITAL ALTERATIONS OF TOOTH DEVELOPMENT

In this first group of dental disorders associated to systemic pathology, mention should be made of taurodontism, characterized by the presence of large pulp chambers that can occupy the entire root. This dental condition is associated to the trichodentoosseous (TDO) syndrome, hypohidronic ectodermal dysplasia and Klinefelter’s syndrome (4). All patients with TDO syndrome present this malformation (5); in contrast, it is only observed in certain hypomaturation variants of amelogenesis imperfecta (6). This marked association between both entities suggests the existence of a genetic determining characteristic referred to as idiopathic dental fluorosis (4). On the other hand, 28.9% of patients with oligodontia suffer taurodontism of one or more, lower molars, with greatly diminished length of the mandibular cuspids and first molars in women (7). In turn, the hypoplastic form of amelogenesis imperfecta can manifest in combination with multiple unerupted teeth, hypercementosis and different root malformation.
ions (8). In animals - specifically, in mice with transgenic cystic fibrosis (9) — anomalies have been observed in the form of soft whitish-blue enamel together with enamel of normal thickness and structure; ameloblasts that rapidly degenerate after the secretory phase, and enamel crystals of granular appearance and low molecular weight.

In the case of odontodysplasia associated to ectodermal dysplasia, clinical manifestations such as hypodontia and hypoplastic enamel appear (10).

Dentinogenesis imperfecta associated to osteogenesis imperfecta constitutes a structural anomaly affecting only the dentine. The teeth appear normal, though their development is abnormal (11); alterations in dentine formation occur in such cases (12). In experimental studies in rats, cyclosporine A has been shown to affect dentine formation, with alterations in the amount of secondary dentine appositioning and the generation of globular dental structures; the pulp is also affected in such situations (13).

**CHROMOSOMAL ANOMALIES**

Turner’s syndrome involves morphological and volumetric dental alterations, with root abnormalities in lower molars and premolars, and reductions in size; the coronal portions of the incisors, canines and premolars are also affected (14), and the mesio-distal diameters are reduced (except in the upper canines) along with the vestibule lingual diameter of some teeth only (15). In Down syndrome the frequency of agenesis is 10 times greater than in the general population, with a higher incidence in males than in females. In order of descending frequency, agenesis affects the maxillary central incisors, the maxillary lateral incisors, the maxillary second premolars, and the mandibular second premolars (16). Microdontia is also observed. Another trisomy-involving chromosome 16, is also associated to dental alterations. In this sense, decreases in the size of different dental organs have been documented in mice, together with the appearance of hypoplasias (17).

In one case of triple X syndrome the congenital absence of teeth was reported, with the presence of only one maxillary central incisor in both the deciduous and permanent dentition (18). Internals with a 45, XO karyotype, reductions in cuspidsurface can be observed, along with decreases in volumen -as reflectedby shortened mesiodistal and vestibulolingual diameters (19). Taurodontism has also been described (20). In Klinefelter’s syndrome (males 47, XXY), important increments in enamel (but not of dentine) have been reported — in contrast to what is seen internally (21). As regards the gnostic condition 45, X/46, XX, 43% of the mandibular premolars have two roots — the proportion being approximately the same as internals with a 45, X karyotype (22).

**RADIATIONS**

Radiotherapy for head and neck cancer produces symptoms such as mucositis, oral dryness and taste alterations (23). A consequence of xerostomia is the increased risk of caries, which in these patients tend to be rapidly evolving, extensive and located in non-habitual zones (24). In children receiving radio-and chemotherapy, the number of dental anomalies has been found to increase (25).

**INTOXICATIONS**

Dental pathology associated to drug ingestion is diverse (26). As regards the production of caries, three groups of drugs can be cited: (a) those containing saccharose as excipient; (b) drugs that depress salivation and there-fore reduce the action of salivary carioprophylactic agents in general (ie., buffer systems, dilution effect, etc.) — including tricyclic antidepressants, antipsychotic drugs, antihistamines, medication for arthritis, analgesics, diuretics, muscle relaxants, antiahythmic drugs, anticonvulsive agents, antidiarreah formulations, antihypertensive drugs, medication for Parkinson’s disease, antispasmodic drugs, anoregeneric agents; and (c) lithium-containing drugs. Drug intoxications can also cause dental discoloration, e.g., topical tin fluoride and systemically administered flours, chlorhexidine (though in this concrete case the dental plaque rather than the actual dental structure is stained), tetracyclines and ciprofloxacin. Regarding morpho-structural alterations of teeth, phenytoin should be mentioned, as it produces shortening, resorption and increased cement depositionturn; local anesthetics are cytotoxic for tooth enamel and can interfere with amelogenesis when introduced under pressure into deciduous tooth ligaments. Additionally, they may cause enamel hypoplasias in permanent dentition. Lastly, doxapram has been reported to induce the pre-mature appearance of dental germs (27).

**GASTROINTESTINAL DISEASES**

One of the most frequent dental alterations is erosion associated to gastrointestinal disorders. An example of this is provided by voluntary regurgitation (28), where the acid gastric contents attack the dental surface, causing progressive dental erosion (wear). In such situations the patient suffers marked dental hard tissue loss in the anterior group, and even in the palatine (lingual) surfaces of the premolars — to the point of exposing the pulp chambers. These alterations may in turn
be associated to dental discoloration. Similar effects are observed in patients with gastro esophageal reflux, where continuous exposure to low pH values leads to irreversible loss of dental substance once the salivary buffering capacity has been overcome (29, 30).

Patients with celiac disease have been found to suffer an increased incidence of amelogenesis imperfecta and other enamel developmental defects (31). In turn, Gardner’s syndrome involves dental anomalies associated to maxillary osteomas.

OSTEODYSTROPHY AND ASSOCIATED CONDITIONS

In two siblings with dwarfism, severe microcephaly has been observed in association with generalized microdontia (32). The appearance of dental dyschromia (gray-yellowish teeth) has also been described, probably as a residue of connective tissue alterations — in one case associated to osteopenia, fetal hydrops and communicating hydrocephalus (33). In three patients in whom retarded growth was associated to hypotonus and hypotrophy, microdontia with extensive diastases were recorded (34).

In the Hallermann-Streiff syndrome, generalized (though sometimes occasional) microdontia can be observed (35), as well as hypodontia, persistent deciduous dentition, enamel developmental defects, late dental development, mandibular hypoplasia, and caries (36). A typical feature of this syndrome is the proximity of the lower molar root apexes to the inferior mandibular margin (37). Ehlers-Danlos mucopolysaccharidosis (type VII syndrome) involves the appearance of microdontia with yellowish discoloration of the teeth, and caries. Radiologically, marked dentinal opacity is observed (38). In turn, Ehlers-Danlos syndrome type I may present imperfect dentinogenesis (particularly of the mandibular incisors), alterations in root size, and occasionally also root hypoplasia or aplasia. Histologically, giant root canals are observed, with pulp calcifications and vascular inclusions (39). In the Winchester syndrome — another example of mucopolysaccharidosis — a clinical case has been reported involving the presence of supernumerary teeth together with unerupted teeth, irregularly spaced teeth and caries (40).

This extensive group of syndromes also comprises the following disorders: infiltrating congenital lipomatosis, with unilateral coronal enlargement or macrodontia, anomalous root formation and chronic periodontitis (41); tumor al calcinosis, where in addition to microdontia and dental structural wear the dental alterations may also involve the root and dental pulp, with the formation of short and bulbous roots, taurodontism of the first molars, pulp calcifications and partial pulp obliteration (42); and pyknody sostosis, where in one documented case sharpened teeth with narrow pulp chambers were observed, together with enamel and root developmental alterations, malocclusion and caries (8).

Dwarfism associated to Grebe chondrodysplasia involves permanent dentition hypodontia along with diminished dental volume; additional findings include delayed formation and eruption of retained deciduous teeth. The jaws are hypoplastic (42). The Russell-Silver syndrome is in turn quite similar as regards the dental manifestations, with hypodontia, microdontia, delayed eruption, an arched palate, and crowding (39, 41); in another case reported in the literature, additional findings comprised the presence of double teeth in the deciduous dentition (40, 41). This form of dwarfism also manifests in the facial region, with an inverted orientation of the labial commissures.

CONCLUSION

The oral cavity is an important, very specific anatomical location with a significant role in many critical physiologic processes, such as digestion, respiration, and speech. It is also unique for the presence of exposed hard tissue surrounded by mucosa. Diseases of the tissues of the oral cavity can be categorized in the various groups: viral, hormonal, fungal, bacterial, dermatological, pharmaceutical, systemic disease, non-cancerous growths, psychiatric disorders, cancer and genetics. The primary and most important factor contributing to oral disease is tobacco use. However, other factors such as: alcohol beverage use, bad oral hygiene, diabetes and other medical conditions affecting the immune system, medications, stress and genetics can all play a role. The mouth is frequently involved in conditions that affect the skin or other multiorgan diseases. In many instances, oral involvement precedes the appearance of other symptoms or lesions at other locations.
Sažetak

**SISTEMSKE BOLESTI KOJE UTIČU NA ZUBNU PATOLOGIJU**

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Prikazan je retrospektivni pregled sistemskih poremećaja koji mogu biti povezani sa zubnom patologijom. Oni su grupisani na sledeći način: (a) kongenitalni dentalni razvojni poremećaji, (b) anomalije hromozoma, (c) zračenja, (d) poremećaji imunskog sistema, (e) trovanja, (f) neurološki poremećaji, (g) gastrointestinalse bolesti, (h) osteodistrofija i slični poremećaji, (i) kožna oboljenja, (j) metabolici poremećaji i endokrini bolesti, (k) kraniofacialni sindromi i udružene kongenitalne malformacije. Za svaki slučaj je ponašob data udružena dentalna patologija.

**Ključne reči:** bolesti zuba, simptomi, sistemskes bolesti, retrospektivni pregled.

**REFERENCES**


