Dental anomalies in fetal alcohol syndrome. A systematic review

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Abstract

Background: Fetal alcohol syndrome (FAS) is the set of signs and symptoms presented by the fetus resulting from maternal alcohol ingestion during pregnancy and preconception periods, such as growth deficit, facial features alterations and delayed neuropsychomotor development, as well as dental anomalies.

Objective: The aim of this systematic review was to investigate the different dental anomalies associated with FAS.

Methods: A Medline search for "fetal alcohol syndrome and dental anomalies" was performed. Further research was conducted for "fetal alcohol syndrome" and its individual oral manifestation. The inclusion criteria were: human clinical studies without a minimum of stipulated patients. Nine studies fulfilled the inclusion criteria and were analyzed accordingly to PRISMA protocol for systematic reviews.

Results: A total of 211 patients with FAS were included in the analyzed studies. The nine studies analyzed included case reports, case series and a review on the subject. Cardiac, respiratory, renal and cognitive defects were the main systemic manifestations, while the main dental abnormalities found were malocclusions, displaced teeth in the dental arch, tooth agenesis and supernumerary teeth.

Conclusion: Although Oral and systemic manifestations are always found in patients with FAS, such common dental anomalies are poorly documented and insufficiently described, and given the importance of such aspects for the patient’s health, we suggest that more studies on this subject are necessary to describe the possible consequences from these anomalies.

Introduction

Fetal alcohol syndrome (FAS) refers to a pattern of congenital malformations seen in children born to women who consumed large amounts of alcohol during pregnancy [1]. FAS hallmarks have been cataloged, quantified, and refined over the years and many investigators have established that most consequences of maternal alcohol consumption during pregnancy are pre and post-natal growth deficiency as well as brain and craniofacial abnormalities [1-8]. FAS was first described in 1973 in a study reporting a variety of growth and developmental abnormalities in 11 children whose mothers drank heavily during their pregnancies [9]. The symptoms can be found from in-utero up to early childhood. There is a wide variety of developmental defects that result from alcohol exposure, including brain abnormalities, central nervous system dysfunctions, and growth deficiencies of developing organs and body systems [5]. These adverse effects on the developing fetus are collectively known as fetal alcohol spectrum disorders (FASD). FAS cause learning, emotion, cognition and motor performance dysfunctions, while also possibly leading to behavioral as well as social issues [10].

Moreover, Orofacial abnormalities are extremely common and an important part of the FAS diagnosis [5,6]. Growth deficits, facial characteristics alterations and neuropsychomotor development delay are all observed [11] and systemic manifestations include low birth weight [8], low weight gain [7], microcephaly (small head) [7], difficulty in learning, language, memory and attention [5], low Intelligence quotient (IQ) [6], changes in vision and hearing [9], difficulties of socialization [10], behavioural disorders [12], delayed cognitive development [11], kidney diseases [8], neurological changes such as seizures, kidney, bone, and congenital heart disease [9]. New-born infants with FAS may be irritable, exhibit muscular hypotonia, severe tremors, and withdrawal symptoms. Mild mental retardation is the most common and serious deficit, while a variety of other anomalies affecting the nervous system, hearing, and organs may accompany FAS [12]. The signs and symptoms are diverse since the passage of ethanol

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Key words: fetal alcohol syndrome, oral health, anodontia

Received: January 20, 2020; Accepted: February 04, 2020; Published: February 13, 2020
through the placenta and the degree of metabolism of ethanol by the maternal liver are both variable [8]. The most common orofacial changes are small eyelid fissures [11], flat facies [12], maxillary hypoplasia [6], short nose [6], long and hypoplastic nasal filter [9], and thin upper lip [10]. The unique facial appearance of FAS patients is the result of changes in 4 areas: short palpebral fissures, flat nasal bridge with an upturned nasal tip, hypoplastic philtrum with a thin upper vermilion border, and a flat midface [11]. Other facial anomalies include micrognathia [13], occasional cleft lip and/or palate and small teeth with defective enamel [3]. Therefore, the aim of this systematic review was to investigate the different dental anomalies described and associated with FAS.

Methods

The present review was carried out in accordance with the PRISMA protocol for systematic reviews [14].

Search strategies: An electronic search in PubMed (MEDLINE), Scopus, Embase, Google Scholar and The Cochrane Library was conducted by two independent reviewers until December 2018, when we researched the following terms following the PICO search strategy (http://askmedline.nlm.nih.gov/ask/pico.php): 'Fetal alcohol syndrome and dental anomalies' [title/abstract] OR 'Fetal alcohol syndrome and dental alterations' [title/abstract] OR 'Fetal alcohol syndrome and dental manifestation' [title/abstract] OR 'Fetal alcohol syndrome and oral manifestation' [title/abstract], OR "Fetal alcohol syndrome and oral findings" [title/abstract] including a literature search strategy.

Inclusion and exclusion criteria: Selection of papers through the inclusion and exclusion of criteria, data extraction, and quality assessment. Meta-analysis was not possible since selected studies did not observe the same variables, methods, participants, and outcomes, which prevented comparisons. A manual search by reading the references of the selected studies was also performed.

Study selection: Prospective or retrospective cohort studies, case series and case reports describing clinical and/or laboratory characteristics of patients with dental anomalies in FAS were included. The titles and abstracts of all reports in the electronic searches were each read by two investigators separately. The studies that fulfilled the inclusion criteria and those that did not present sufficient information in the title or in the abstract, were fully assessed. A cross-check assessment of the articles was performed by another author to guarantee that the proper studies were selected according to the inclusion and exclusion criteria.

Data extraction: We independently extracted study data through specific extraction forms using Microsoft Excel software. For the study selected, the following information was extracted (when available): year and country of publication, article type, number of cases, sex, if consanguinity existed in the patients (which could lead to a search), systemic manifestations and dental alterations described by the authors.

Analysis: All the patients’ clinical information and diagnostic criteria were evaluated for data descriptive cross-referencing, considering the information given by the authors, wherein the main dental alterations were observed in FAS-affected patients.

Results

The searches returned a total of 74 records. After the duplicates were removed, 35 records remained (Figure 1). All 74 articles were selected through the titles and abstracts and 29 records were excluded based on the exclusion criteria. Of the excluded records, 65 did not focus on the presence of dental anomalies in the FAS, of which 4 were reports of animal tests, 47 reported facial changes without anomalies mentions and 11 were deemed irrelevant to the topic, resulting in 9 papers that met the search criteria, classified as follows: 3 case series, 3 case reports, 2 original research article and 1 letter to the editor [15-24]. All cited and selected papers had in their methodology and in their discussion the focus on showing oral, facial and systemic alterations in patients with FAS (Table 1). All articles followed the 1966 IOM (Institute of Medicine Diagnostic) criteria, even those that were published before that date were reviewed. The paper by Streissguth et al., [4,5] was the first article describing malocclusions and changes in dental position found in patients with FAS [16] while also describing other facial and systemic alterations of patients with FAS. However, Chu et al., was the first one that specifically described the number and shape of dental anomalies in serial cases of 22 patients and cited the presence of agenesis, twisting teeth and diastema. 8 of the papers cited the presence of oral clefts in the reported cases [15-23]. In the paper by Andrade [24], it was clearly shown the presence of dental alterations and abnormalities and made their correlation with systemic renal alterations, which are derived from the same genetic defect of numeral dental alterations.

Discussion

FAS is characterized by a pattern of congenital anomalies that include pre and/or postnatal growth retardation, central nervous system involvement with neurologic abnormalities, developmental delay or intellectual impairment, and various facial birth defects [1-5,8,15-18,22]. It has been reported that during the prepubertal age, FAS patients have a characteristic emaciated appearance, which may not remain a feature in adolescence, at least in females [10-12,23]. Diagnostic criteria

Figure 1. Flowchart of process of systematic literature search, using PRISMA flow.
<table>
<thead>
<tr>
<th>Author</th>
<th>Location</th>
<th>Article type</th>
<th>Sex</th>
<th>Number of patients</th>
<th>Consanguinity</th>
<th>Systemic alterations</th>
<th>Orofacial alterations</th>
<th>Dental anomalies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Streissguth et al. [5]</td>
<td>USA</td>
<td>Serial Case</td>
<td>7 (F) 4 (M)</td>
<td>11</td>
<td>Yes</td>
<td>Borderline intelligence, eustachian tube dysfunction, Cardiac anomalies</td>
<td>Midface hypoplasia, oral cleft, ptosis, protrusion and maxillary and mandibular retraction</td>
<td>Malocclusions</td>
</tr>
<tr>
<td>Jackson &amp; Hussain [19]</td>
<td>USA</td>
<td>Letter to the editor</td>
<td>Not described</td>
<td>6</td>
<td>Yes</td>
<td>No descrip</td>
<td>Facial Dysmorphism, left lip and palate, maxillary hypoplasia, short ram of mandible</td>
<td>Delay eruption</td>
</tr>
<tr>
<td>Pećina- Hrncević &amp; Buljan [23]</td>
<td>Croatia</td>
<td>Case report</td>
<td>3 (F) 3</td>
<td>Yes</td>
<td>-</td>
<td>Dermatoglyphic alterations, and cutaneous syndactyly and clinodactyly</td>
<td>Microcephaly, hypertelorism, malformed and low-positioned auricles, cleft palate, aplasia of the uvula, wide root of the nose, saddle nose, enamel hypoplasia</td>
<td>Enamel hypoplasia,</td>
</tr>
<tr>
<td>Webb et al. [17]</td>
<td>USA</td>
<td>Case report</td>
<td>1 (F) 1</td>
<td>Yes</td>
<td>-</td>
<td>Strabismus</td>
<td>Short palpebral fissures, Low, broad root of nose, Upturned nose, Short nose-upper lip distance, Flattened midface profile</td>
<td>Malocclusion, Retromandibular mandible</td>
</tr>
<tr>
<td>Church et al. [22]</td>
<td>USA</td>
<td>Serial Cases</td>
<td>14 (F) 8 (M)</td>
<td>22</td>
<td>Yes</td>
<td>Hearing problems</td>
<td>Oral Clefts, microcephaly, malformed Noses, (small, upturned, cleft, and flat nasal bridge), deformed ears (narrow canals, prominent/deformed pinnae, and otosclerosis), deformities about the eyes (epicanthal folds, hypertelorism, ptosis, and narrow palpebra Fissures), malformations to the soft tissue around the mouth (thin upper lip, flat philtrum, and downturned mouth)</td>
<td>Malocclusion at various levels, Absent teeth, displaced or Rotated teeth, diastema</td>
</tr>
<tr>
<td>Autti-Ramo et al [18]</td>
<td>Finland</td>
<td>Serial Case</td>
<td>76 (F) 1 (M)</td>
<td>77</td>
<td>Yes</td>
<td>Systemic alterations in: Cardiac, Genitourinary, Skeletal, Gastrointestinal kidney, Pulmonary, Vision, Strabismus</td>
<td>Occipitofrontal circumference, Palpebral fissure length, Attention deficit/hyperactivity, Fine motor dysfunction, Midfacial hypoplasia, “Railroad Track” ears, Proptosis, Epicanthal folds (non-racial), Flat nasal bridge, Antverted nare, Long philtrum, This vermilion border of upper lip, Prognathism</td>
<td>Dental crowding</td>
</tr>
<tr>
<td>Santa’ana and Tosello [20]</td>
<td>Brazil</td>
<td>Research</td>
<td>-</td>
<td>Yes</td>
<td>-</td>
<td>Cardiac malformations, Ventricular septal defects, pulmonary artery hypoplasia and interruption of aortic arch, type A. Other less common abnormalities in skeletal, urinary, ocular, immune and renal system</td>
<td>Cleft palate, cleft lip, And dentition anomalies malformed noses, (small, upturned, cleft, and flat nasal bridge) and deformed ears (narrow canals, prominent/deformed pinnae, and otosclerosis)</td>
<td>Crowded incisors, maxillary overjet, and open bite, Malocclusion, Enamel Hypomaturated, Agenesia, dental crowding, Malformation and arching Of the maxilla and failure In exfoliation and eruption</td>
</tr>
<tr>
<td>Naidoo et al. [21]</td>
<td>South Africa</td>
<td>Research</td>
<td>45 (F) 45 (M)</td>
<td>90</td>
<td>Yes</td>
<td>Immature Skeletal</td>
<td>Oral cleft, orofacial malformation</td>
<td>Enamel Hypomaturated, Agenesia, dental crowding, Malformation and arching Of the maxilla and failure In exfoliation and eruption</td>
</tr>
</tbody>
</table>

**Table 1.** Papers used in the systematic review of fetal alcohol syndrome, showing the main oral and systemic manifestations of the syndrome.
for FAS developed by IOM simplified the identification of classic FAS cases. When the phenotype is less than classic or atypical, confirming the diagnosis becomes difficult, if not impossible [17,18,20]. Early diagnosis is essential for appropriate intervention on children affected by prenatal alcohol exposure and can reduce their risk of facing social difficulties later in life as well as employment issues due to behavioural and learning difficulties or even lawbreaking occurrences as a result of impulsive behaviour and lack of inhibition [15-24]. In the papers presented, FAS diagnosis was made through the association of facial and systemic alterations, and the presence of several dental anomalies, such as agenesis, malocclusions, diastemas, twisting tooth and dental crowds [15-24]. Naidoo et al observed a reduced age of dental maturation in 90 patients with FAS compared to a normal patient showing a slight alteration in development characterized by delayed enamel maturation [21]. Andrade et al. [24] observed its applicability in the case report in which boy shows enamel hypomaturation and agenesis of several teeth due to a acute kidney disease, which is also described in the work of Autti-Ramo et al. (2005) and Sann’tanna et al. (2006).

It has been documented that alcohol consumption during pregnancy can be a cause of orofacial cleft [25]. Munger et al., [26] reported that maternal alcohol consumption during pregnancy was associated with an increased risk of cleft lip with or without cleft palate. Furthermore, the offspring of alcohol-consuming mothers showed a higher chance of developing cleft lip and palate were higher compared to women who did not consume alcohol during pregnancy [26]. The number of distinct teratogenic mechanisms works simultaneously along different biochemical pathways and at different physical locations in the developing embryo [27]. The ways in which this alcohol-induced mechanism damage the fetus depends on several variables, such as time, frequency, and the amount of maternal alcohol consumption during pregnancy; the state of health and habits of the mother as well as the genetic constitution of the mother and the fetus. These teratogenic effects directly affect dental anomalies presence in FAS [28]. Dental and maxillofacial alterations such as orofacial fissures are due to the teratogenic alcohol effect on the fetus, which as described in Sant’anna’s et al. (2006), causes alteration in the MSX1 and MSX2 gene, both participating in the maxillary process’s fusion odontogenic formation, a hypothesis reinforced by Naidoo et al., [13,21]. The most frequently occurring anomalies were crowded incisors, maxillary overjet, and open bite [16-18,20,21]. Levels of plaque and gingival bleeding were high, which was not an unexpected finding since these children come from backgrounds in which oral hygiene is not a high priority [19].

Conclusion

In this systematic review, where the papers published in the main research platforms were scanned, it is observed that there are few studies that explain and show the direct relationship between oral manifestations and SAF, evidencing the various dental anomalies that these patients present such as agenesis, malocclusions, diastemas, twisting tooth and dental crowds. More studies are needed to explore these dental alterations, since they cause aesthetic and functional problems in these patients who already have several isthmian alterations, thus making the effects of SAF in these patients more discreet.

Sources of funding

None

Conflict of interest

None

Prospero register

CRD42019133863

Acknowledgment

The Minas Gerais State Research Foundation-FAPEMIG, Minas Gerais, Brazil and the National Council for Scientific and Technological Development - CNPq, Brazil.

Authors’ contribution

RSA designed and wrote the manuscript, RMP and JAQ reviewed the manuscript, DRBM assisted in the search methodology, HMJ oriented and reviewed the manuscript.

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