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Abstract: The Cornelia de Lange syndrome (CdLS) is a genetic disorder that affects newborns between 1/10.000 and 1/100.000, and whose diagnosis is established based on medical history and clinical examination, characterized by growth retardation and development, hirsutism, structural abnormalities in the limbs and distinctive facial features. The aim of this paper is to provide a systematic review of the literature on oral manifestations of this rare disease, describing 7 published cases of CdLS in the indexed literature (PubMed/MEDLINE) - the first corresponds to the year 1980 and the last in 2009. Dental problems among these patients are common and include: delayed eruption, malocclusion, micrognathia, dental malpositions, extensive caries or periodontal disease. In any case, the delayed eruption, which may be the cause of diastemas and/or malocclusion is the most common oral finding associated with CdLS - which early diagnosis is key.

Keywords: Cornelia de Lange syndrome, oral manifestations, systematic review.

I. Introduction

The Cornelia de Lange syndrome (CdLS) is a rare developmental malformation characterized by a learning disability by a variable degree of intellectual impairment, short stature due to their growth retardation, abnormalities in the limbs (oligodactyly, amputations...) and a particular facial dysmorphism [1]. They may also have other malformations in organs such as the heart or kidney [2]. It was first described by Dr. Cornelia de Lange, who published two cases of children with similar traits [3]. The syndrome has also been called Brachmann de Lange, as was Dr. Brachmann in 1916 who described similar symptoms in one patient [4]. The estimated incidence varies from 1/10.000 to 1/100.000 live births [5]. Virtually all cases are sporadic but may occur following an autosomal dominant pattern [6]. The NIPBL gene, one of the 3 known involved, is mutated in 50% of cases [7]. However, the diagnosis continues to be established with the clinical history and examination. Distinctive facial findings are widely accepted as diagnostic criteria of CdLS (Table 1) [8]; curved eyebrows, confluent and well-defined, long and curly eyelashes, anteverted nostrils and mouth downturned with a thin upper lip, giving it the aspect of crescent shaped mouth. However, cases have been described with atypical facial features that can cause uncertain diagnosis [9]. There is no specific treatment for CdLS but therapies aimed at improving behavior, stimulation and neurodevelopmental program along with a number of drugs to control seizures that these patients can undergo [10].

Oral manifestations of CdLS have been briefly described in the medical literature. As for the dental literature, there are some articles describing the oral pathology of these patients. The present study aims to provide a systematic review of the literature on oral manifestations of this rare disease.

II. Material And Methods

A literature search on the subject in question on the basis of PubMed/MEDLINE data up to May 10th, 2016 was performed the following keywords were used: Cornelia de Lange syndrome AND OR dental oral manifestation. The articles were independently assessed by two reviewers (JGS, VMPR). The inclusion criteria were: articles in English generated by the MEDLINE database, submit (at least) a case with oral manifestations and to describe oral characteristics of the disorder and the treatment used in this case. Of the 58 articles found, only six met the requirements.
III. Results

The first article published in the dental literature is that of Scully in 1980 [11], who described two clinical cases (median age 17.5 years). The first one was a caucasian male 18 years of age, who underwent a dental prophylaxis with an ultrasonic device for presenting chronic marginal gingivitis due to poor oral hygiene. He did not need any other dental treatment afterwards. He was given oral hygiene instructions that later showed a significant improvement of his periodontal health. Meanwhile, the second case was a 17 year-old black male who underwent a dental prophylaxis with an ultrasonic device to treat chronic periodontitis also due to poor oral hygiene. He received oral hygiene instructions as well. He died a year later as a result of pneumonia.

We did not find any other case until 1993. It was Barrett et al [12] who described the case of a 22 year-old female who presented dental pain in several mandibular teeth that also were affected by caries. The patient was epileptic and was treated with clonazepam (2mg twice/day) and carbamazepine (200mg daily). She went under general anesthesia for multiple surgical extractions (26, 34, 36, 45 and 46). A mucoperiosteal flap was raised, and an ostectomy performed due to the extreme bone density and large roots. The patient was sutured and there were no postoperative incidents. She required intravenous administration of factor VIII:C on the morning of the surgery as well as the next morning because she had a variant of von Willebrand disease.

In 2005, Gupta et al [13] presented the case of an 11 year-old male who was in need of orthodontic treatment for having upper front teeth in an extreme buccal position. Two years later, in 2007, Grau Carbó et al [14] reported a case of a 29 year-old female who had temporal teeth, as well as, several ectopic molars extracted. She also received the appropriate periodontal treatment to have dental health. All the procedures were performed under general anesthesia. And seven days later, she was evaluated again to check her oral health, and educate her family about the importance of daily oral hygiene procedures. It was recommended that the patient come to check-ups every 4 months to prevent further disturbances.

Guadagni et al [15] in 2008 described the case of a 3 year-old male who went for a dental check-up due to the neuropsychiatrist’s concern that the child was experiencing dental pain, which could be increasing the frequency of the seizures. The child’s parents received oral hygiene instructions and dietary advice. The patient was monitored every 2 months to assess the level of hygiene, growth and development of the jaws and tooth eruption. A systemic treatment with fluoride was administered as well as topically. Orthodontic treatment was discarded for lack of patient cooperation, and an absence of chewing and swallowing reflex activity.

Finally, in 2009, Toker et al [16] presented a case of a 10 year-old male that was experiencing dental pain, and whose mother reported having a history of preterm births. The patient had the affected teeth extracted under general anesthesia and there were no complications during the process or in the postoperative period.

Of the studies published to date regarding CdLS, we found 5 articles that included a single case report [12-16] and an article that included a sample of two patients [11]. (Table 2). The mean age of the patients described in this review of 5 clinical cases and a series of 2 cases was almost 16 years (exactly 15.7 years), the age range being between 3 and 29 years. Of the revised dental literature, only 7 people with this rare syndrome were subjected to an oral examination, being too small a sample to draw firm conclusions as to the degree of oral pathology and the problems that could arise while managing them in the dental clinic.

IV. Discussion

There are oral manifestations derived mainly from the medication used for the treatment of CdLS [12]. These patients are difficult to evaluate because of a communication problem, the lack of language and the altered pain threshold, which could mask the presence of pain [15].

Just as anxiolytics and anticonvulsants can cause xerostomia, stomatitis, glossitis, erythema multiforme, dysgeusia, etc., it needs to be considered the possible interactions of these drugs, as in the case of acetylsalicylic acid or erythromycin [12]. Therefore, a detailed medical history, containing all the drugs used by the patient to control the disease is extremely important in order to avoid undesirable side effects.

Thus, we can only distinguish a group of manifestations derived from the specific conditions of CdLS. Orofacial manifestations in patients with CdLS found in the literature are described in Table 2. Their assessment indicates that they are not exclusive of the syndrome, since they can also present in other diseases characterized by the presence of oral patterns, seizures or difficulties with a proper oral hygiene. However, there are some features, such as delayed eruption [11,13,16], anterior open bite [12,15] or gingivitis [11,14], which are more common among patients with CdLS. The pathology associated with poor hygiene, like an early-onset [16] or multiple form of tooth decay [12,14] and periodontal disease [11,14], is closely related to this syndrome due to cognitive difficulties they present.

The unique presence of upper central incisors shaped like those of Hutchinson’s suggests a possible link between the symptoms of CdLS and Hutchinson syndrome [16]. A controlled diet, topical fluoride application and periodic check-ups are crucial in the management of these patients [11,14,15] and may limit the use of sedation and general anesthesia for the most extreme cases. In terms of behavior management, it is necessary to consider limitations in the communicative ability of the patient, mental retardation and lack of
attention. The techniques should always be tailored to each individual case. Some authors have used general anesthesia [12,14,16] due to possible health risks, and the lack of cooperation necessary for proper dental treatment. It can also be exploited to perform other procedures as in the case presented by Grau Carbó et al [14] who made a dilated esophagus with fibrogastroscopy.

V. Figures And Tables

Pubmed/MEDLINE search without date restriction until May 10\textsuperscript{th}, 2016:
(Cornelia de Lange syndrome) AND ((dental OR oral) AND (manifestations OR findings))

<table>
<thead>
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<th>No</th>
<th>Main criteria</th>
<th>With</th>
<th>Secondary criteria</th>
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<tbody>
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<td>Facial</td>
<td></td>
<td>Synophrys (arched, fine eyebrows)</td>
<td>And ( \geq 3 )</td>
<td>Long eyelashes: Short nose, anteverted nares; Long, prominent philtrum; Broad or depressed nasal bridge; Small or square chin; Thin lips, down-turned corners; High palate: Widely spaced or absent teeth</td>
</tr>
<tr>
<td>Growth</td>
<td>( \geq 2 )</td>
<td>Weight below 5th centile for age; Height or length below 5th centile for age; OFC below 2nd centile for age</td>
<td>Developmental delays or mental retardation; Learning disabilities</td>
<td></td>
</tr>
<tr>
<td>Development</td>
<td>( \geq 1 )</td>
<td>Developmental delays or mental retardation; Learning disabilities</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Behavior</td>
<td>( \geq 2 )</td>
<td>Attention deficit disorder ± hyperactivity; Obsessive-compulsive characteristics; Anxiety Constant roaming; Aggression; Self-injurious behaviour: Extreme shyness or withdrawal; Autistic-like features</td>
<td></td>
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</tr>
<tr>
<td>Musculoskeletal or</td>
<td></td>
<td>Reduction defects with absent forearms Small hands and/or feet (below 3\textsuperscript{rd} centile) or Oligodactyly</td>
<td>Only and ( \geq 2 ) and ( \geq 3 )</td>
<td>5th finger clinodactyly; Abnormal palmar crease: Radial head dislocation/abnormal elbow extension; Short 1st metacarpal/proximally placed thumb Bunion; Partial 2,3 syndactyly toes; Scoliosis: Pectus excavatum; Hip dislocation or dysplasia</td>
</tr>
<tr>
<td>or</td>
<td>None of these</td>
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Figure 1. Flow diagram of the literature search.
There are no acknowledgements.

References


Table 1. Diagnostic criteria for CdLS (Kline et al, 2007) [8].

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<tr>
<th>Authors</th>
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<th>Age (years)</th>
<th>Oral manifestations</th>
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Table 2. Oral manifestations of Cornelia de Lange syndrome found in the dental literature.

VI. Conclusion

In conclusion, this review shows that oral habits should be strongly considered in these patients because it may lead to malocclusion and anterior open bite or an arched palate. However, the most representative clinical feature of CdLS is a delayed eruption, which may be the cause of other events such as the presence of diastemas and/or malocclusions that are difficult to treat without early diagnosis, due to the lack of collaboration of these patients.

Acknowledgements

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