Solitary median maxillary central incisor syndrome: Case report

Eduardo Machado*, Patricia Machado**, Betina Grehs***, Renésio Armindo Grehs****

Abstract

**Introduction:** The presence of a single median maxillary central incisor is an uncommon event in the population. The prevalence of the Solitary Median Maxillary Central Incisor (SMMCI) syndrome is about 1:50,000 live births, occurring more in women. This alteration in the development of the dental occlusion is characterized by structural malformations, over all in midline region of the patient. The early diagnosis and the adequate treatment of this syndrome are of great importance, therefore this condition can be an indication that the patient can present other severe congenital malformations, not having to consider the SMMCI a simple dental anomaly. The orthodontic procedures, in these cases, vary depending on the degree of involvement of bone structures of the maxilla, the occlusion in itself, and mainly of the midpalatal suture. **Objectives:** To discuss, based on scientific evidence, important aspects related to the SMMCI and present a clinical case of female patient with SMMCI, which was submitted to orthodontic treatment in the Children’s Dental Integrated Clinic of the Federal University of Santa Maria - RS/Brazil. **Conclusion:** According to the critical analysis of literature, it is very important to correctly early diagnose this condition, since there is the possibility of this syndrome to be associated with other problems of development. Moreover, the patients affected by SMMCI should be attended by a multidisciplinary health team in order to optimize the clinical results and recover the quality of life of these patients.

**Keywords:** Solitary median maxillary central incisor. Single median maxillary central incisor. SMMCI. Orthodontics.
INTRODUCTION
The congenital absence of upper central incisors is a rare condition, while the presence of a single central incisor also is an uncommon event.21 The prevalence of the Solitary Median Maxillary Central Incisor (SMMCI) syndrome, also known as Single Median Maxillary Central Incisor Syndrome, occurs in 1:50,000 live births, with higher involvement of women. In this syndrome, developmental defects occur due to unknown factors operating in utero about the 35th–38th day from conception and are characterized by structural malformations, mainly midline defects in the patients.11,24

Thus, the purpose of this study is to discuss, within a context based on scientific evidence, and illustrate, with a case report, relevant aspects concerning this condition.

CASE REPORT
A eight years and three months old patient, female, Caucasian, Brazilian, presented to the Children’s Dental Integrated Clinic at Federal University of Santa Maria/RS (Brazil) for evaluation. After the initial clinical examination, the patient was selected and referred to the division of Orthodontics at the Children’s Dental Clinic. Once accepted at the Division of Orthodontics of this clinic, the patient was well attended, her clinical history and the records of physical-clinical examinations were obtained and the orthodontic records necessary for diagnosis and treatment planning were requested. During clinical examination a very significant alteration was observed, the presence of a single central incisor, compatible with SMMCI, and maxillary atresia, as shown in figures 1, 2 and 3.

Regarding the presence of systemic changes,
the patient’s parent reported no involvement. This evaluation is important, since SMMCI may be associated with other developmental problems such as congenital nasal abnormalities,\textsuperscript{1,4,11,15,16,17} growth deficiencies,\textsuperscript{8,22} holoprosencephaly,\textsuperscript{6,28} format changes and craniofacial morphology,\textsuperscript{25} congenital heart disease,\textsuperscript{8,10} among other local and systemic changes. However, there are studies that found no relationship between SMMCI and systemic changes.\textsuperscript{5,27}

Some authors also found associations between SMMCI and SHH gene mutations\textsuperscript{9,10,12,14,19,20} and deletions in parts of chromosome 18p\textsuperscript{2,7} and/or of chromosome 7q.\textsuperscript{10,18,26} Thus, an evaluation of a geneticist can find some association between SMMCI and chromosomal abnormalities.

The orthodontic treatment plan comprised a Phase I, which consisted of rapid maxillary expansion (RME), as well as support and interaction with Prosthetic Dentistry, Pediatric Dentistry and Oral and Maxillofacial Surgery specialties. At the end a Phase II was scheduled with fixed orthodontic treatment. Furthermore, the patient was referred to a multidisciplinary health team, including pediatricians, geneticists, speech therapists and psychologists, since this anomaly may be associated with other developmental problems.

**DISCUSSION**

The involvement of SMMCI was initially reported by Scott\textsuperscript{23} who described a girl with the presence of a solitary median maxillary central incisor, as an isolated finding. Another case of SMMCI was verified by Fulstow,\textsuperscript{8} but the patient showed apart from the single central incisor, short stature, congenital heart disease, microcephaly and scoliosis. Some factors that may be associated with SMMCI are the pituitary gland dysfunction and short stature, whereas in a study involving patients with SMMCI, 7 subjects had short stature and 5 were deficient in growth hormone production.\textsuperscript{22} However, Wesley et al\textsuperscript{27} reported two cases of SMMCI in subjects with normal stature, while Cho and Drummond\textsuperscript{5} reported three cases of SMMCI in three Chinese girls with no growth deficiencies or systemic involvement.

According to DiBiase and Cobourne,\textsuperscript{6} the most common cause of a missing maxillary

---

FIGURE 3 - A) Periapical radiograph, which confirms the presence of a solitary maxillary central incisor. B) Occlusal radiograph, confirming the presence of a solitary maxillary central incisor.
central incisor is trauma, or more rarely hypodontia. When dental absence has no explanation in the patient’s clinical history, a genetic analysis can show results. It is important to recognize the SMMCI when in an unknown etiology, because it may indicate a risk factor for holoprosencephaly. Thus, the role of the orthodontist is extremely important in the diagnosis of this condition, which must refer the patient for genetic testing to investigate other possible developmental disorders.

The SMMCI may be associated with various congenital nasal anomalies such as choanal atresia, intra-nasal stenosis and nasal pyriform aperture stenosis. Choanal atresia consists in a bone or membranous obstruction of the posterior nasal aperture caused by a failure in the oronasal disintegration. The intra-nasal stenosis is a bony narrowing of the nasal cavity between the pyriform aperture and the posterior choanae, whereas the nasal pyriform aperture stenosis is an anterior nasal obstruction secondary to the bone growth of the nasal processes of the maxilla. It is important to note that the clinical aspects of the above changes are similar, and often a computed tomography is required for definitive diagnosis, being that prenatal diagnosis of SMMCI can be done through magnetic resonance imaging.

Thus, several studies have looked at the association of nasal obstructions and SMMCI. Arlis and Ward evaluated six patients with congenital stenosis of the nasal pyriform aperture and found that of these, 4 had SMMCI. Lo et al found in their results that 63% of patients with congenital stenosis in the nasal pyriform aperture also presented SMMCI, while Hall et al found that among 21 patients with SMMCI, all had a positive relationship with a history of nasal congenital obstruction, whereas choanal atresia and intra-nasal stenosis were found in 7 and 8 patients respectively. Already, Levison et al reported two cases of neonatal children with nasal obstruction due to stenosis of the choanae, which had an association with single maxillary central incisor, a fact verified by computed tomography.

The presence of chromosomal defects was observed in some children who had SMMCI. Dolan et al found chromosomal abnormalities in children with a single central incisor, with deletion of parts of chromosome 18 (18p), which was also reported by Aughton et al. Nonetheless, Masuno et al reported deletion in the terminal portion of chromosome 7q, which was also found by Hall and Tubbs and Oakes. Another factor that seems to be associated with the SMMCI is a mutation of the SHH gene.

For Yassin and El-Tal, the appearance of a solitary incisor in place of the two central incisors may occur due to fusion of two neighboring teeth or to agenesis of a tooth germ. However, this can be associated with other systemic disorders such as autosomal dominant holoprosencephaly, growth retardation and midline developmental defects. Becktor et al evaluated the intermaxillary suture, the eruption pattern of the single central incisor and growth of the maxilla in a group of patients with SMMCI. The sample consisted of 11 patients with SMMCI, who underwent orthopantomographs, dental and lateral cephalometric radiographs. The X-rays showed that the intermaxillary suture was abnormal anterior to the incisive foramen, however, the horizontal and vertical growth of the maxilla was normal.

Analyzing 10 patients (9 girls and 1 boy) aged between 8 and 17 years who presented SMMCI, Kjaer et al examined the clinical characteristics and craniofacial morphology of this group of patients. Intra and extra-oral photographs, profile radiographs, orthopantomograms and study casts were analyzed. The study results showed that the craniofacial morphology of nine girls with SMMCI compared with normal craniofacial
parameters, showed a short anterior cranial base, a short, retrognathic and posteriorly inclined maxilla, and a retrognathic and posteriorly inclined mandible, and morphological changes in the sella turcica were found in five patients examined. Moreover, this group of patients had characteristics such as: nasal obstruction, septal deviation, absence of the frenum of the upper lip, and a complete or incomplete mid-palatal ridge. Thus, the presence of SMMCI should not be considered as a simple dental anomaly, because it may be associated with other clinical characteristics and craniofacial malformations.

Tabatabaie et al25 evaluated the neurocranial and craniofacial morphology of children with SMMCI using profile radiographs and cephalometric analysis. The sample comprised 13 children (12 girls and 1 boy) aged between 7 and 17 years. Cephalometric evaluations were compared with standard measures. The study results showed that the size of the neurocranium, the maxillary prognathism and inclination, the mandibular prognathism and inclination of lower incisors are significantly decreased in patients with SMMCI. But, the mandibular inclination, vertical jaw relationship and mandibular angle are significantly increased in patients affected by SMMCI. The data from this study showed that the occurrence of SMMCI is a sign of anomaly development, associated with deviations in neurocranial size and shape and in craniofacial morphology.

According to Hall,10 the etiology of SMMCI is uncertain and may be associated with mutations in SHH gene (I111F) in chromosome 7q, with a positive correlation with congenital nasal malformations. These teeth erupt and develop in the midline of the maxillary arch, both in primary and permanent dentitions. The presence of SMMCI may be associated with some common congenital abnormalities such as moderate to severe intellectual disability, congenital heart disease, cleft lip and/or palate and less frequently, microcephaly, hypopituitarism, strabismus, duodenal atresia, scoliosis, hypothyroidism, absent kidney, micropenis and ambiguous genitalia. Short stature can be found in children. The diagnosis of SMMCI should be performed at 8 months of age, but can be done at birth and possibly prenatal, between the 18th and 22nd week of gestation by ultrasound examination. In patients with SMMCI rehabilitation should be undertaken in accordance with the anomalies presented by individuals: choanal stenosis requires surgical treatment, short stature should be approached with growth hormone therapy, and the presence of single maxillary central incisor should be a requirement for an multidisciplinary treatment involving the specialties of Orthodontics, Prosthetic Dentistry and Oral Surgery.

Cho and Drummond5 suggest that early diagnosis of SMMCI is extremely important, because it is a sign that the patient may present with other severe congenital malformations. If they are pediatric patients they should be seen together with the pediatrician. In three patients evaluated by these authors,5 all were female and had no growth deficiencies or any systemic involvement. The dental management consisted in preventive and orthodontic approaches, and in two cases expansion of the upper arch was performed, moving the solitary central incisor to one side and obtaining space for placement of osseointegrated implant or prosthesis on the other side.

CONCLUSIONS

This case report has a great clinical importance under the viewpoint of the orthodontic treatment necessary to solve this dental occlusion anomaly. The simple fact of a malocclusion being present, associated to a maxillary atresia, synthesize the functional severity that this type of case represents, requiring adequate oral rehabilitation, as well as an integral attention to the health of patients suffering from SMMCI.
Dental procedures for patients with SMMCI vary with the degree of commitment that it causes. Orthodontic procedures are extremely important for the return of function and aesthetics to the patient, requiring an interdisciplinary approach with other dental specialties for optimizing clinical outcomes. Moreover, it is important that the patient should be attended by a multidisciplinary health team, including pediatricians and other medical professionals, geneticists, speech therapists and psychologists, since this anomaly may be associated with other developmental problems and systemic changes.


Contact address
Eduardo Machado
Rua Francisco Trevisan, nº 20, Bairro Nossa Sra. de Lourdes
CEP: 97.050-230 - Santa Maria / RS, Brazil
E-mail: machado.rs@bol.com.br

Submitted: August 2008
Revised and accepted: October 2008