Rubinstein-Taybi syndrome: Dental manifestations and management

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Rubinstein-Taybi syndrome (RSTS) is an uncommon genetic disorder characterised by a typical facies, small stature, broad angulated thumbs and intellectual impairment. Dental changes are a minor, yet significant component of the condition. Craniofacial growth retardation in RSTS is frequently complicated by unerupted teeth, while dental caries is related to the inherent intellectual deficit. Dental problems necessitate interdisciplinary management in terms of oral surgery, conservative dentistry, periodontics and orthodontics. When affected individuals are unco-operative, certain dental procedures may warrant general anaesthesia. In these instances, dental and medical staff will combine their expertise to enhance the well-being of the patient. In addition, specific dental changes may alert the medical practitioner to the possible diagnosis of RSTS. In this article we document the oro-dental manifestations and review the oro-dental approach in the management of three patients with RSTS. Our experience in South Africa may be relevant to other countries at a similar stage of development.

The three patients reported in this article were assessed by a multidisciplinary dental team from the University of the Western Cape (UWC), Bellville, Cape Town, and medical genetics staff from UCT.

Case 1
An affected white girl born in 1985 was first seen at the UWC-UCT Dental Genetic Clinic in 2003. Her mother reported an uneventful pregnancy followed by an emergency caesarean section because of fetal distress. Her non-consanguineous parents and her elder brother were unaffected.

The girl was ‘floppy’ during infancy and her developmental milestones were delayed. Her intellectual compromise persisted during childhood, and special educational facilities were necessary. She wore a bib until the age of 5 years because of constant drooling. Enuresis persisted until puberty. In adolescence, her joints were hypermobile, and ligamentous laxity and instability of the left knee necessitated orthopaedic intervention. An ophthalmological operation was undertaken in 2002 to rectify deviation of her right eye.

On examination at the age of 18 years, she had the characteristic facies of RSTS, notably a prominent forehead, microcephaly, down-slanting palpebral fissures, epicanthal folds, strabismus, a broad nasal bridge, a pear-shaped nose with the nasal septum extending below the alae, and a small oral opening (Fig. 1). Her hands appeared normal, but both big toes (halluces) were broad.

Intra-oral examination revealed a high-vaulted palate, and several teeth were absent from the oral cavity. Her malocclusion (Fig. 2) was characterised by edge-to-edge occlusion, a bilateral posterior cross-bite, a bilateral posterior open bite, and lack of posterior tooth support on the right side.

A pantomogram showed the clinically missing permanent teeth retained in the alveolar bone (Fig. 3). Analysis of the available jaw space failed to reveal any discrepancies that would affect the eruption of...
these teeth. Cephalometric analysis revealed a retrognathic mandible and maxilla with a high mandibular plane angle and cantered occlusal plane. The maxillary incisors had an angulation and inclination within the normal range, while the mandibular incisors were retroclined and retruded.

Routine chromosomal analysis did not reveal any abnormality. Molecular techniques, including fluorescence in situ hybridisation (FISH) testing and sequencing of the CREBBP gene, can be used to confirm the clinical diagnosis of RSTS, but are currently not available in South Africa.

Case 2
A South African black girl born in 2003 was diagnosed as having RSTS at 20 months of age and followed up regularly at Red Cross War Memorial Children’s Hospital, Cape Town, until the age of 4 years. Her family history was negative.

She was born at term following a pregnancy complicated by hyperemesis gravidarum. She had no significant medical illnesses, but her developmental milestones were severely delayed: she started walking at 2 years of age and spoke her first words at 3 years of age.

Her growth was retarded with height below the third centile and weight on the third centile. She was microcephalic and had typical facial features of RSTS, including down-sloping palpebral fissures, a beaked nose with the columella extending below the alae nasae, a prominent lower lip and a grimacing appearance of the mouth. The palate was high-arched. The terminal phalanges of all fingers were broad, while the thumbs and halluces were strikingly broad. She had a slight pectus excavatum and both thumbs and halluces were broad. No other family member was affected.

During the first visit, mother and child were given instruction in oral hygiene and their diet was analysed. The boy was not co-operative in the dental chair, but it was possible to apply topical fluoride to the teeth. Regular follow-up visits indicated that his oral health status had improved significantly. A finger brush was suggested for cleaning the maxillary teeth, as the gums in this area were very sensitive. Arrangements were made for conservative treatment and extractions of carious teeth under general anaesthesia.

The boy returned to the dental clinic 4 years later when he was in the mixed dentition stage. Apart from poor oral hygiene and a few cavities, no additional dental anomalies were noted. The sequence of tooth eruption was within the normal range for his age. His extreme anxiety prevented any radiographs from being obtained.

Discussion
In 1957, Michail et al.[1] described a 7-year-old boy with the distinctive features of RSTS in a French orthopaedic journal. In 1963, unaware of the aforementioned report, Rubinstein and Taybi[2] documented the syndromic association of broad thumbs and toes with facial abnormalities, thereby delineating the condition that now bears their conjoined eponym.

A mutation in the CREBBP gene (OMIM 600140) at the chromosomal locus 16p13.3
is detectable by the FISH technique in about 15% of persons with RSTS (RSTS1). With sequence analysis, this figure rises to 30-50%. In a minority of individuals, the EP300 gene (OMIM 602700) at 22q13 has been implicated (RSTS2). Considerable intragenic heterogeneity has been recognised in both forms of RSTS. In general, the craniofacial abnormalities and intellectual impairment are more severe in RSTS1 than in RSTS2.

The severity of the manifestations of RSTS is very variable, and the case reports presented in this article provide a perspective of the range of involvement. It is relevant that management, including dental measures, is determined by the severity, or otherwise of the feature in question. The main practical problem in RSTS is mental retardation, and affected persons are normally incapable of independent existence and adequate self-care. Guidelines for the general surveillance and medical management of the RSTS have been proposed by Wiley et al. 10-12

Considerations pertaining to dental management include the use of general anaesthesia if cardiac or respiratory abnormalities are present. Possible challenges during intubation result from anterior positioning and mechanical inadequacies or weakening of the larynx, and there is a risk of aspiration pneumonia. Cardiac abnormalities and arrhythmias have been reported in RSTS, 10,11 and if present, require evaluation before conservative or invasive dental treatment. Skeletal and soft-tissue changes in the head and neck region resulting in upper-airway obstruction can lead to mouth breathing and recurrent pneumonia.

The oral features of RSTS have been reviewed extensively by Hennekam and Van Doorne 6,10 and Bloch-Zupan et al. 11,12 Dental talon cusps are present in a high proportion of persons with RSTS, 11,12 but this feature was absent in our patients. All three had high-arched palates – often associated with malocclusion and tooth malalignment. Hypo- and hyperdontia may also occur. Caries resulting from inadequate dental hygiene is frequent, and in older children may be the result of the intellectual disability and motor impairment that characterise the syndrome. Problems with feeding during early development could be the cause of the nursing caries in cases 2 and 3.

Optimal oral and dental health is of the utmost importance in the general well-being of a patient with RSTS. Good oral health enables problem-free assimilation and digestion of food, which translates to a physically healthy individual. Caries and periodontal disease affect quality of life. If individuals with RSTS cannot articulate coherently, they may suffer from dental pain for weeks or months. In turn, this may affect their eating habits, resulting in malnutrition and an impaired immune response. Bleeding gums may prevent caregivers or parents from continuing routine oral hygiene practices, and together with halitosis may result in social isolation and poor self-esteem.

Proposed management protocols

The dental treatment plan that we proposed for patient 1 encompassed 6-monthly follow-up visits for monitoring the eruption of the remaining teeth. As she showed no signs of a space discrepancy, surgical exposure and assisted eruption would have been unwarranted. Subsequent radiographs showed evidence of eruption with open root apices. Maxillary arch expansion for the correction of the bilateral posterior cross-bite was undertaken. In addition, the anterior teeth were levelled and aligned, where necessary, to facilitate the eruption of the canine and premolar teeth in their appropriate positions. A fixed lingual arch-wire was placed in the mandibular arch to maintain the space for the erupting canine and premolar teeth.

In view of her age and the developmental status of her dentition, patient 2 would ideally have been recalled at 6-monthly intervals to enable adequate monitoring of her growth and development. Management of patient 3 was difficult because of the severity of his intellectual impairment. Extractions and conservative treatment were undertaken under general anaesthesia without complications. The boy returned to the dental clinic every 6 months for cleaning and prophylaxis.

Conclusion

The oro-dental complications of RSTS are important but neglected aspects of the disorder. On the basis of our experience in Cape Town it has become apparent that the multifaceted problems necessitate a team approach towards dental management and possible anaesthesia. Equally, in an environment where facilities may be suboptimal, multidisciplinary collaboration is crucial for effective management and for the well-being of the affected person.

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References