Dental Management in a Patient with Turner Syndrome with Dental Anomalies: A Case Report

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Abstract

Turner syndrome (TS) is a chromosomal disorder caused by monosomy of the X chromosome, with complete or partial absence of the second sex chromosome. Anomalies of root morphology have been found to occur more often in patients with TS, which make endodontic treatment challenging and requires special handling. The patients with TS may also have systematic problems such as cardiac or renal malformations, so in treating these patients it is important for clinicians not only to be aware of the characteristic intraoral findings, but also to make the patients have regular dental check-ups to prevent oral complications in advance.

An 12-year-old girl who had been diagnosed with TS at the age of 10 years was referred due to discomfort in the bilateral mandibular premolar regions. Dens evaginatus and taurodontism were detected in all the mandibular premolars characteristically. The bilateral mandibular first premolars had three roots and the bilateral mandibular second premolars had periapical lesion with two roots. Due to the complexity of the root canal anatomy, root canal treatment were completed with a dental microscope to ensure adequate visualization. After 2 years of regular follow-up examinations, there were no clinical sign or symptom associated with the teeth, and no periapical lesion, was found.

This case report describes the characteristic oral features and dental management of TS patients.

Key words: Turner syndrome, Dental anomalies, Taurodontism

I. Introduction

Turner syndrome (TS) is a chromosomal disorder caused by monosomy of the X chromosome, with complete or partial absence of the second sex chromosome[1]. The estimated prevalence of TS is approximately 1 in 2,500 live female births[2]. The syndrome has some typical characteristics: short stature, hormonal disorders such as estrogen deficiency; and cardiac or renal malformations[3]. TS has been associated with aortic coarctation and dissection, and thoracic vascular anomalies occur in 50% of cases[4].

The most common oral findings in patients with TS are small teeth, short roots, mandibular sagittal deficiency, and a high palatal arch[5]. Abnormal mandibular premolar root numbers have also been observed[6,7].

This study reports on the dental treatment of mandibular premolars in patient with TS, with dens evaginatus and taurodontism, which is known to be associated with abnormalities
in the sex chromosomes[8]. It was performed with the aid of a dental microscope due to the complex root canal anatomy and anomalies of root morphology that is often seen in patients with TS.

II. Case report

An 12-year-old girl was referred from a local dental clinic to the Department of Pediatric Dentistry, complaining of discomfort in the bilateral mandibular premolar regions. Endodontic treatment of both mandibular second premolars had been initiated 2 days previously in a local dental clinic, and the root canals were temporarily filled with a root canal filling material (Fig. 1).

The patient had been diagnosed with TS at the age of 10 years. At the first visit, the patient's height and weight were 122.5 cm and 28.4 kg, respectively, and they were below the third percentile according to the Korean Children and Adolescents Growth Standards.

The patient exhibited abnormally low posterior hairline which is characteristically found in patients with TS (Fig. 2).

The patient underwent balloon dilatation at the age of 1.5 years due to coarctation of the aorta, a congenital heart disease. At the age of 10.5 years, the patient received growth hormone therapy to improve the short stature associated with TS, and underwent strabismus surgery under general anesthesia.

Oral examination showed the hypomineralized type of generalized enamel hypoplasia, with opaque lesions and brown discoloration, in the entire erupted dentition (Fig. 3). Dens evaginatus was detected in all premolars, but not in the bilateral mandibular second premolars where endodontic treatment had been undergoing. The previous crown shape of the bilateral mandibular second premolars, which were pain sites, could not be confirmed as the treatment had been initiated already. However, given that acute pain occurred suddenly on both sides without caries, the pain was presumed to be due to the fracture of the tubercle of dens evaginatus.

Fig. 1. Initial Panoramic radiograph.

Fig. 2. Extraoral lateral photograph. The patient exhibited low and webbed neck and abnormally low posterior hairline which is characteristically found in patients with TS.

Fig. 3. Initial intraoral photographs. Hypomineralized type of generalized enamel hypoplasia were observed with opaque lesions and brown discoloration, in the entire erupted dentition.
Radiographic examination showed that the maxillary right first and second premolars, the maxillary left second premolar, and the mandibular left and right second premolars had two roots; the maxillary left first premolar and the bilateral mandibular first premolars had three roots (Fig. 4). All the mandibular premolars showed elongated pulp chambers and apical displacement of pulpal floor, suggesting taurodontism.

Clinical and radiological examinations revealed periapical abscesses with sinuses in the bilateral mandibular second premolars, and we decided to perform root canal treatment. Due to the cardiac history of the patient, we consulted the pediatrician about precautions before endodontic treatment.

Selective grinding and pit and fissure sealing were scheduled for the other premolars with dens evaginatus, and after 4 days a flowable, light-cured composite resin was layered incrementally on the tubercles and surrounding surfaces.

About 2 weeks after the patient's first visit, the patient took 1500 mg of amoxicillin for prophylaxis 1 hour before root canal treatment as the pediatrician had recommended the use of prophylactic antibiotics before each treatment. Root canal washing and Vitapex® (Neo Dental Chemical Products Co. Tokyo, Japan) removal were performed on the mandibular bilateral second premolars. Due to taurodontism, the root canal orifices were difficult to identify and the root canal was C-shaped. We thus decided to operate under a dental microscope (Global Surgical, St. Louis, MO) to ensure adequate visualization. The mandibular bilateral second premolars were confirmed to have two roots and three root canals, which were mesiobuccal, mesiodistal and lingual, respectively. After Vitapex removal, the root canals were measured and instrumented with ProTaper (Dentsply Maillefer, Ballaigues, Switzerland) NiTi instruments. But the treatment had to be delayed for 2 months since the patient was scheduled to undergo strabismus surgery again. During that period, root canals were filled with calcium hydroxide paste.

At the patient's next visit, the root canals were instrumented with a ProTaper NiTi file, with canal formation and dilatation. The symptoms were confirmed to have disappeared and the canals were filled with gutta percha (GP) cones. After root canal filling, the upper core was formed with composite resin (Fig. 5, 6).

One month after root canal filling, clinical and radiological examinations were performed to confirm the prognosis of the treatment site. Periodic screening was performed every 3 months thereafter (Fig. 7). At the 2-year follow-up examination, no clinical sign or symptom associated with the teeth, and no periapical lesion, was found on panoramic radiographs (Fig. 8).

![Fig. 4. Initial periapical views. The maxillary premolars exhibited dens evaginatus. Taurodontic root morphology and complicated root canals were observed in all mandibular premolars. (A) Maxillary right premolars, (B) Maxillary left premolars, (D) Mandibular right premolars.](image1)

![Fig. 5. Endodontic treatment procedures of mandibular right second premolar. (A) Initial periapical view, (B) Working length determination, (C) Master cone adaption, (D) Filling of three canals.](image2)
TS is a chromosomal disorder characterized by complete or partial absence of an X chromosome in females. On karyotype examination, approximately 50% of patients show complete loss of the X chromosome and the remaining patients show a multitude of chromosomal abnormalities, including partial loss of the X chromosome and mosaicism[9].

Most patients with TS have characteristic clinical features, such as short stature, delayed pubertal growth spurt, primary amenorrhea, and gonadal dysgenesis[10]. They may also have visual or hearing abnormalities, and/or an elevated risk of thyroiditis and other autoimmune diseases[11]. Approximately 30% of patients with TS have congenital heart defects[12-15]. Therefore, when establishing dental treatment plans for patients with TS, dentists should consider the need for prophylactic antibiotics to prevent infective endocarditis and, in some cases, cardiological consultation. The development of intelligence is normal in most patients with TS, and these patients have no unusual difficulty with behavioral control.

In the case reported here, the patient visited the Department of Pediatrics with the chief complaint of short stature. The patient had been diagnosed with TS at the age of 10 years and had received growth hormone therapy since that time. Height and weight of the patient were below the third percentiles, according to the Standard Growth Curve for Korean Children and Adolescents. As a result of the balloon dilatation procedure that the patient underwent at the age of 1.5 years to treat coarctation of the aorta, the patient had to take preventive antibiotics before invasive procedures. As the patient's intellectual
level was normal, and the patient was very adaptive to treatment; no special behavioral control was necessary[16].

The facial features of TS are a senile (frequently heart-shaped) face; multiple ocular findings, depressed corners of the mouth; prominent, low-set ears[10]; and a short, broad, webbed neck[17,18]. In this case, the patient exhibited several features consistent with TS, such as a short and webbed neck and low posterior hairline (Fig. 3).

Reduced tooth crown size and altered tooth morphology have been observed in patients with this syndrome[5,19,20]. Many authors have explained that the X chromosome controls tooth size and shape, as well as tooth root morphology[21,22]. Enamel hypoplasia is observed frequently in patients with TS. Abnormalities of the X chromosome affect the secretion of enamel, thinning it, as demonstrated quantitatively, and causing enamel dysplasia, as observed qualitatively[23]. In the case reported here, intraoral findings included opaque white lesions and brown discoloration in the entire erupted dentition, caused by the hypomineralized type of generalized enamel hypoplasia.

Anomalies of root morphology also have been found to occur more often in patients with TS than in control subjects[23]. Increased numbers of first- and second-premolar roots have also been observed[24-27]. The most frequently observed deformities are two-rooted mandibular first and second premolars[26,27]. Kusiak et al.[24] examined the morphology of the mandibular premolar roots in 40 females with TS; they reported that two-rooted mandibular first premolars were observed in 34% of cases in the X monosomy (45,X) group, and two-rooted mandibular second premolars were observed in 39% of cases in the monosomy group. On the other hand, approximately 98% of the mandibular first premolars were single rooted; the incidence of two roots was 1.8% and three roots were found in 0.2% of all teeth studied. In anatomic studies, almost all (99.6%) mandibular second premolars were found to be single rooted. The occurrence of two roots (0.3%) and three roots (0.1%) is extremely rare[28,29]. Thus, it is extraordinary that all the mandibular premolars of this patient had two or three roots.

Furthermore, considering that the general prevalence of taurodonts was reported to range 2.5 - 11.3% of the human population, it is assumed to be a very rare case that all the mandibular premolars were diagnosed to be taurodonts[30]. Varrela et al.[31] suggested that the X chromosome influencing development of enamel may also be involved in the development of taurodontism. Taurodontism also has been reported that it is sometimes associated with several syndromes and anomalies, including amelogenesis imperfecta, ectodermal disturbance, Down syndrome, Klinefelter’s syndrome, and others[31,32].

As the canal orifices of the bilateral mandibular second premolar roots were difficult to navigate, we decided to use a dental microscope during root canal treatment to ensure adequate visualization and treatment completion. It was also reported that careful exploration of the grooves between all orifices, especially with magnification had been recommended, since each taurodont tooth may have extraordinary root canals in terms of shape and number[33].

Patients with TS have an elevated risk of caries due to weak and thin enamel structure[21]. In addition, complicated root structures in the premolar region render root canal treatment difficult in these patients. And awareness and consideration of endodontic treatment difficulties in patients with TS is a crucial clinical factor.

In the treatment of TS patients, dentist should be aware of the possibility of systemic complications and strive for periodic dental follow-up for preventive maintenance. And if endodontic treatment is needed, the dentist should also take account for the complexity of the root shape and the variation of the root number in these patients.

IV. Summary

The patient in this case had TS and showed dental anomalies, such as dens evaginatus and taurodontism in all the mandibular premolars. Anomalies of root morphology have been found to occur more often in patients with TS, which make endodontic treatment challenging and require special handling. Due to the complexity of the root canal anatomy of the mandibular premolars of the patient, root canal treatment were completed with a dental microscope to ensure adequate visualization. Due to cardiac history, the patient had to take prophylactic antibiotics before each treatment.

The patients with TS may also have systematic problems such as cardiac or renal malformations, so in treating these patients it is important for clinicians not only to be aware of the characteristic intraoral findings, but also to make the patients have regular dental check-ups to prevent oral complications in advance.
References

국문초록

치아형태이상을 가진 터너 증후군 환자의 치과적 관리

이한이 전공의 · 신세영 전문의 · 김재곤 교수 · 이대우 교수 · 양연미 교수

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터너 증후군이란 한 개의 X 염색체와 전부 혹은 부분이 결여된 X 염색체를 나타내는 염색체 이상 질환이다. 치근 형태 이상이 터너 증후군 환자에서 정상에서보다 호발한다고 보고되었는데, 치근의 이러한 특징은 근관치료를 까다롭게 만들고, 특수한 처치를 요하기도 한다. 또한 터너 증후군 환자들은 종종 심장 또는 신장의 기형과 같은 전신적인 문제를 동반하기도 하므로, 임상의는 터너 증후군의 특징적인 구내 소견을 인지하고, 사전에 구내 합병증을 예방하기 위해 환자들이 정기적으로 치과 검진을 받을 수 있게 해야 한다.

10세 때에 터너증후군 진단을 받은 12세 여자 환자가 하악 양측 소구치 부위의 불편함을 주소로 내원했다. 특징적으로 하악의 모든 소구치에서 우상치와 치외치가 관찰되었다. 하악 양측 제1소구치는 세 개의 치근을, 근단부 병소를 보이는 하악 양측 제2소구치는 두 개의 치근을 가지고 있었고. 근관의 복잡한 해부학적 특징 때문에 치과 현미경을 이용한 적절한 시야를 확보하여 근관치료를 마무리했다. 2년간의 정기적인 관찰 결과 해당치아와 관련된 임상적 징후나 증상, 치근단 병소는 관찰되지 않았다.
본 증례 보고를 통해 터너 증후군 환자의 구강내 특징과 이의 치과적 관리에 대해 보고하고자 하였다.