Dental Considerations in Ehlers-Danlos Syndrome: A Case Report

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Introduction: This study aimed to elaborate a case with several complications and recommended some suggestions for dental management of these individuals.

Case Presentation: Ehlers-Danlos syndrome (EDS) is a relatively rare condition involving a group of patients with inherited connective tissue disorders. Main manifestations of this condition are fragile and hyperextensible skin as well as joint mobility. A 6.5 year-old-girl referred with clinical presentation of frequent unexplained bruises leading to the characteristic clinical features of ED syndrome. Gentle and atraumatic approach in every way is the key in medical and dental management of such patients as minimum pressure could still lead to severe bruises even during a simple dental examination.

Conclusions: All clinical findings were directly related to the features described for EDS. There are occasions when dentists would be the first health professionals to see these individuals at a young age; knowing the classic signs of EDS could help in identifying and managing them properly.

Keywords: Ehlers-Danlos Syndrome; Oral; Dental; Manifestations; Children; Management

1. Introduction

Ehlers-Danlos syndrome (EDS) is a relatively rare condition involving a group of patients with inherited connective tissue disorder. This syndrome is caused by 3 fundamental mechanisms, including deficiencies of collagen-processing enzymes, dominant negative effects of mutant collagen, chains, and haploinsufficiency. The main findings of EDS cases include articular hypermobility, skin extensibility, and tissue fragility (1-3). The prevalence of EDS has been reported as 1 in 5000 to 100000 live births in different communities; however, the epidemiology of the specific types is largely unknown. EDS affects both sexes and has no racial predisposition (4-6). EDS has a wide range of expressing pattern depending on the type of collagen being affected. Villefranche introduced a classification system for EDS, which contained 6 major types. Variations were mainly based on the clinical, biochemical, and molecular differences (2, 3) (Table 1). The classical type characterized with hypermobility is known to be the most common form of EDS having type V collagen deficiency in patients with classic EDS, which demonstrates marked skin hyperextensibility, wide atrophic scarring, and significant joint hypermobility (1). Other findings comprised smooth velvety skin, molluscoid pseudotumors over pressure points, subcutaneous spheroids on the forearms or chins, muscle hypotonia, ecchymosis, and tissue fragility (5). Surgical or traumatic bruising can vary from mild to moderate and can be accompanied by delicate hyperextensible skin (7). Repeated dislocation of TMJ could be reported along with epicanthus, strabismus, narrow nasal bridge, and shaggy hair in facial area (8).

Intraoral manifestations include highly fragile mucosa and frequent periodontal tissue injuries. This is usually seen following minor oral surgeries such as a simple tooth extraction (9). Bleeding tendency is higher in cases of EDS compared to normal cases. Early-onset of generalized periodontitis has been reported in some cases (10, 11). The tongue is very soft with Gorlin sign being visible in almost 50% of the patients with EDS (9). The palate is usually deep and dome-shaped (9, 12). Dental anomalies include enamel hypoplasia, some degrees of root deformity, pulp stones, missing and supernumerary teeth (8, 13-16). This article aimed to focus on various clinical complications of EDS with more emphasis on oral and dental findings and management of a 6.5 year-old-female with EDS.
Table 1. Different Types of EDS With Varying Degrees of Tissue Involvements

| Villefranche Types | Collagen Defects | Main Features |
|--------------------|-----------------|---------------|
| Classical          | Type V          | Skin hyperextensibility, wide atrophic scarring, significant joint hyper mobility |
| Hypermobility      | Unknown         | Generalized joint hyper mobility and musculoskeletal pains |
| Vascular           | Type III        | Ecchymoses, smooth, muscle wall fragility, small joint hyperextensibility, translucent skin, easy and excessive bruising |
| Kyphoscoliosis     | Lysyl hydroxylase | Hypotonia, joint laxity, congenital scoliosis, ocular fragility |
| Arthrochalasia     | Type I          | Severe generalized joint hypermobility with recurrent subluxations, congenital bilateral hip dislocation |
| Dermato-sparaxis   | Procollagen N-peptidase | Skin fragility leading to sever ecchymoses |

2. Case Presentation

A 6.5-year-old Iranian female was presented to the pediatric dental clinic at Shahid Beheshti University of Medical Sciences in September 2010. Patient had a familial history of EDS with parents being cousins. Patient was reported to have been born with asphyxia after full term pregnancy, which led to a period of incubation for 11 days. A history of urinary tract infections (UTI) was also noted with moderate degrees of kidney problem (hydronephrosis) and a prolonged respiratory problem. Several different drugs are administered to tackle organ problems, including salbutamol inhalation aerosol (Ltd co., China) and the cap of pamidronate (Bioadvantex Pharma, USA). An informed consent form was signed by parents for the GA (general anesthesia) dental procedure and the report preparation. Complete Blood Count (CBC) was evaluated with other vital signs as a routine part of the case investigation. CBC test results indicated that all measures were within the normal range.

Extra oral examination revealed severe hyperelasticity of the skin in various parts (Figure 1). Hypermobility of the joints was noted (Figure 2), which was detected soon after birth. Therefore, the patient had to be controlled for several joint dislocations while skin examination showed slight bruises and dystrophic scars. Intraoral examination revealed no dental abnormality in shape or number of the teeth as expected. However, dental caries was present with poor oral hygiene (Figure 3 A) with an unusual tongue posture (Figure 3 B). These findings were further confirmed by an orthopantomograph view (Figure 4). For treatment and management of the case, several teeth were extracted with the rest being restored in one visit under general anesthesia (Figure 5).

3. Discussion

Widely known cases of EDS suffer from varying degrees of extra movements and these clinical signs play an important role in identifying their specific type. Further laboratory investigations were performed on skin biopsy with results showing disturbed or abnormal collagen structure in classical type (5).
Major diagnostic criteria for classical type include skin hyperextensibility, wide atrophic scars, and significant joint hypermobility. Similar clinical signs were observed in the presented case. Several other reported diagnostic criteria include recurrent joint dislocation, positive family history, bleeding tendency, prolapsed mitral or tricuspid valves (5). This patient manifested all 3 major criteria as well as recurrent joint dislocations and positive family history. Absence of any musculoskeletal pain is considered as a major sign in hypermobile type of EDS, also the absence of ecchymosis and or scoliosis. Interestingly, patients with EDS are reported to carry several dental anomalies such as hypoplasia of enamel and dentin with short dilacerated roots, calcified pulp (pulp stone), congenital missing, and even presence of supernumerary teeth. However, the present case was free from any dental anomalies with only caries as the major dental problem (8, 13-16). Several active caries associated with hypoplastic enamel may somehow been related to the collagen defects associated with EDS (17).

Based on the age and the number of caries teeth, it is recommended to treat such children under general anesthesia as there is a high risk of temporomandibular joint (TMJ) dislocation in cases seen on chairside. No complication has been reported for dental treatment under general anesthesia from intubation, medication, or dental works. Overall, cases with EDS d, require pre-operative cardiac consultation and antibiotic prophylaxis cover prior to any major dental operations due to the presence of the mitral valve prolapsed (4).

Pillsbury showed that fragility of skin and blood vessels could lead to scar and hematoma formation following...
any traumatic event in EDS patients (18). Due to the possibility of complications following any treatment within the oral cavity, care should be taken as even low traumatic forces could lead to severe bleeding. Excessive gingival hemorrhage is reported as a common finding following teeth brushing (5, 8, 16). Surprisingly, such finding was not reported in the present case. Prolonged and repeated bleeding is reported following teeth extractions (5, 19, 20). It is critical to control laboratory blood test results, including prothrombin time (PT), partial thromboplastin time (PTT), and international normalized ratio blood test (INR) as a routine part of evaluation in patients with EDS. Blood count elements were found to be in normal range for the present case. Teeth were planned to be treated in one visit under general anesthesia. Care should be taken when suturing the socket walls of an extracted tooth in order to ensure a minimally traumatizing procedure for treatment under general anesthesia and to reduce the risk of extra trauma from local anesthetic administration. Reasonable care should be implemented when suturing the extraction sites in such individuals as any excess pressure and tension could further damage the already fragile and elastic tissues (16, 19, 21). An acrylic removable appliance is suggested to be used as a replacement for stitches to cover the surgical site (16, 21, 22). Inferior alveolar nerve block injection is banned because of its high risk of tissue damage (8, 23). Surgical flaps involving mucoperiosteom need maximum care as it is associated with high risk of tissue collapses. TMJ hypermobility and high dislocation should also be considered when undergoing dental treatment in patients with EDS. It is also recommended to arrange appointments in short periods. No orthodontic treatment is recommended to be scheduled until the full eruption of the first molars.

All initial orthodontic procedures are best provided along with routine dental treatments under general anesthesia in a single visit course for those suffering from EDS. As collagen defects influence periodontal ligament (PDL) and gingival fibers, light orthodontic forces are suggested along with a longer retention period (24). Higher frequency of root resorption and soft tissue defects are among the problems reported to be associated with orthodontic treatment of patients with EDS. Jone reported a high risk of mucosal ulceration related to bracket positioning in a case of EDS (25). Pourshahidi et al. had spotted similar findings in another similar case under dental observation (26). Fragility of gingival tissue and blood vessels could make the performance of any routine dental procedure highly sensitive. Therefore, special care should be taken when there are any interties for small dental works for chair side circumstances of patients with EDS.

The diagnosis of this case was EDS with the typical defects of collagen structure reflected in different tissue damages, including the tongue, fingers and toes. Medical and dental professionals are encouraged to see these patients willingly but carefully in order to deliver atraumatic, gentle care with a more preventive approach.

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Authors’ Contributions
All authors have been equally contributed in preparation of various parts of the article.

References
1. Mao R, Bristow J. The Ehlers-Danlos syndrome: on beyond collagen. J Clin Invest. 2008;110(9):2053–9.
2. Beighton P, De Paepe A, Steinmann B, Tsipouras P, Wenstrup RJ. Ehlers-Danlos syndromes: revised nosology, Villefranche, 1997. Ehlers-Danlos National Foundation (USA) and Ehlers-Danlos Support Group (UK). Am J Med Genet. 1998;77(1):31–7.
3. Yassin OM, Rihani FB. Multiple developmental dental anomalies and hypermobility type Ehlers-Danlos syndrome. J Clin Pediatr Dent. 2006;30(4):337–41.
4. Abel MD, Carrasco LR. Ehlers-Danlos syndrome: classifications, oral manifestations, and dental considerations. Oral Surg Oral Med Oral Pathol Oral Radiol Endod. 2006;102(5):582–90.
5. Steinmann B, Royce PM, Superici-Furga A. The Ehlers-Danlos Syndrome: Connective tissue and its heritable disorders: molecular, genetic, and medical aspects: New York: Wiley-Louisie; 2002.
6. Sollecito TP, Sullivan KE, Pinto A, Stewart J, Korostoff J. Systemic conditions associated with periodontitis in childhood and adolescence. A review of diagnostic possibilities. Med Oral Patol Oral Cir Bucal. 2005;10(2):342–50.
7. De Paepe A, Mallat F. Bleeding and bruising in patients with Ehlers-Danlos syndrome and other collagen vascular disorders. Br J Haematol. 2004;127(5):491–500.
8. Letourneau Y, Perusse R, Buthieu H. Oral manifestations of Ehlers-Danlos syndrome. J Can Dent Assoc. 2000;67(6):330–4.
9. Piette E, Douiniala R. [Juvenile parodontolysis symptomatic of Ehlers-Danlos syndrome, a sporadic case?]. Acta Stomatol Belg. 1980;77(3):217–29.
10. Welbury RR. Ehlers-Danlos syndrome: historical review, report of two cases in one family and treatment needs. ASDC J Dent Child. 1989;56(3):220–4.
11. Hartsfield KJ, Kousseff BG. Phenotypic overlap of Ehlers-Danlos syndrome types IV and VII. Am J Med Genet. 1990;37(4):465–70.
12. Fridrich KL, Fridrich HH, Kempf KK, Moline DO. Dental implications in Ehlers-Danlos syndrome. Oral Surg Oral Med Oral Pathol. 1990;69(4):431–5.
13. Norton LA, Assael LA. Orthodontic and temporomandibular joint considerations in treatment of patients with Ehlers-Danlos syndrome. Am J Orthod Dentofacial Orthop. 1997;111(1):75–84.
14. Barabas GM. The Ehlers-Danlos syndrome. Abnormalities of the enamel, dentine, cementum and the dental pulp: an histological examination of 13 teeth from 6 patients. Br Dent J. 1969;126(1):509–15.
15. Hoff M. Dental manifestations in Ehlers-Danlos syndrome. Report of a case. Oral Surg Oral Med Oral Pathol. 1977;44(6):664–71.
16. Gosney MB. Unusual presentation of a case of Ehlers-Danlos syndrome. Br Dent J. 1987;163(2):54–6.
17. Iltingberg G, Hagberg C, Noren JG, Nietzsche S. Aspects on dental hard tissues in primary teeth from patients with Ehlers-Danlos syndrome. Int J Paediatr Dent. 2009;19(4):282–90.
18. Karrer S, Landthaler M, Schnalz G. Ehlers-Danlos type VIII. Review of the literature. Clin Oral Invest. 2000;4(2):66–9.
19. Barabas GM, Barabas AP. The Ehlers-Danlos syndrome. A report of the oral and haematological findings in nine cases. Br Dent J. 1967;123(10):473–9.
20. Recant BS, Lipman JS. The Ehlers-Danlos syndrome. A case requiring oral surgery. *Oral Surg Oral Med Oral Pathol*. 1969;28(4):460–3.

21. Hughes CL. Odontectomy in treatment of Ehlers-Danlos syndrome: report of case. *J Oral Surg*. 1970;28(8):612–4.

22. Sadeghi EM, Ostertag PR, Eslami A. Oral manifestations of Ehlers-Danlos syndrome: report of case. *J Am Dent Assoc*. 1989;118(2):287–91.

23. Sacks H, Zelig D, Schabes G. Recurrent temporomandibular joint subluxation and facial ecchymosis leading to diagnosis of Ehlers-Danlos syndrome: report of surgical management and review of the literature. *J Oral Maxillofac Surg*. 1990;48(6):641–7.

24. Norton LA. Orthodontic tooth movement response in Ehlers-Danlos syndrome: report of case. *J Am Dent Assoc*. 1984;109(2):259–62.

25. Jones ML. Orthodontic treatment in Ehlers-Danlos syndrome. *Br J Orthod*. 1984;11(3):158–62.

26. Pourshahidi S, Ebrahimii H, Taghavi Zenouz A, Andisheh Tadbir A. Oral Manifestations of Ehlers-Danlos Syndrome and Presentation of a Case. *Iran Red Crescent Med J*. 2009;11(2):206–9.