Congenital Adrenal Hyperplasia: A Case Report With Premature Teeth Exfoliation and Bone Resorption

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Abstract

Congenital adrenal hyperplasia (CAH) is an inherited autosomal recessive disorder characterized by insufficient production of cortisol. The aim of this case report was to present a child with CAH, premature exfoliation of primary teeth and accelerated eruption of his permanent teeth related to bone resorption. A 4.5-year-old Caucasian boy with CAH and long-term administration of glucocorticoids was referred for dental restoration. Clinical examination revealed primary molars with worn stainless steel crowns, severe attrition of the upper canines, and absence of the upper incisors. Before the completion of treatment, abnormal mobility of the first upper primary molars and the lower incisors was detected, and a few days later the teeth exfoliated prematurely. Histologic examination revealed normal tooth structure. Alkaline phosphatase and blood cells values were normal. Eruption of the permanent dentition was also accelerated. Tooth mobility was noticed in the permanent teeth as soon as they erupted, along with bone destruction. Examination revealed an elevated level of receptor activator of nuclear factor-κB ligand and lower-than-normal osteoprotegerin and vitamin D levels. The patient was treated with vitamin D supplements, and his teeth have been stable ever since. CAH is a serious chronic disorder appearing in children with accelerated dental development and possibly premature loss of primary teeth.

Congenital adrenal hyperplasia (CAH) is an inherited disorder that leads to insufficient production of cortisol caused by abnormality in 1 of the 5 enzymes that are necessary for its biosynthesis. Glucocorticoid and mineralocorticoid administration is the therapy of choice. However, chronic administration of corticosteroids may cause growth retardation, drug-induced osteoporosis, glucose intolerance, and infection susceptibility.

Increased bone resorption in patients with CAH has been noted in the past and has been related to alterations in receptor activator of nuclear factor-κB ligand (RANKL) and osteoprotegerin (OPG). This effect might be caused by the long-term administration of corticosteroids or the excessive secretion of adrenocorticotropic hormone. Also, bone resorption has been related with loss of teeth in patients with osteolytic diseases.

In addition, adrenal hyperactivation has been found to increase infection susceptibility and increase the risk for development of periodontal disease. Periodontal disease is related to loss of teeth.

Accelerated tooth eruption has been reported in patients with CAH. Long-term therapy with glucocorticoids and excessive hormonal secretion have been associated with the acceleration of dental development. Recently, bone resorption has been found to be correlated with accelerated eruption of teeth in mice.
Premature exfoliation of teeth has not been reported in patients with CAH. However, susceptibility to periodontal disease and bone resorption can be the cause for primary exfoliation of primary teeth and accelerated eruption of its successors.

The aim of this case report is to present a child with CAH with premature exfoliation of primary teeth and accelerated eruption of his permanent teeth caused by bone resorption.

**CASE REPORT**

A 4.5-year-old white boy was referred in June 2010 to the postgraduate clinic of pediatric dentistry of the Dental School, University of Athens, for dental restoration. The patient had a BMI of 13.5, which is below the 5th percentile for boys of that age. Medical history included preterm birth with normal karyotype (46 XY). Medical history included asthma controlled with medication and history of pneumonia at the age of 9 months. The child had been hospitalized several times for viral infections and surgical procedures (e.g., hypospadias, cryptorchism, and hydrocele). The main medical condition was congenital hyperplasia of adrenal cortex, treated with systemic intake of hydrocortisone (9 mg/m² per day) and potassium levothyroxine (T4, 5 μg/kg) daily. Dental history included uncooperative behavior with the dentist, toothbrushing once a day by the mother, and severe bruxism during the day and night.

Clinical examination revealed the presence of worn primary dentition and absence of the maxillary primary incisors. Parents reported extraction of the upper incisors due to abscess caused by pulp exposure after severe attrition. Patient had no signs of caries or enamel hypomineralization.

The treatment plan included an individualized preventive program and rehabilitation of dentition with stainless steel crowns applied under general anesthesia (GA). Dental treatment under GA was a necessity not only because the child was uncooperative but also because patients with CAH need administration of supplemental doses of glucocorticoids to cope with dental anxiety.

Before the scheduled appointment for dental treatment under GA, the patient presented for emergency care because of gingival swelling, halitosis, and abnormal mobility of first upper primary molars. The patient received a triple daily dose of supplemental steroids, and the teeth were extracted under local anesthesia. In addition, abnormal mobility of the primary lower central incisor was recorded, and radiographic examination revealed severe bone loss (Fig 1). The incisor fell off during the examination. All teeth were sent for histologic examination to define the cause of the multiple tooth loss, and laboratory blood tests were requested.

Regarding premature exfoliation of primary teeth, the patient was checked for aggressive periodontitis, immunodeficiency (neutropenia, leukocyte adhesion deficiency, leukemia), hypophosphatasia, Langerhans histiocytosis, Papillon–Lefèvre syndrome, diabetes mellitus, hyperthyroidism, dentinal dysplasia, acatalasia, Singleton–Merten syndrome, Hajdu–Cheney syndrome, Coffin–Lowry syndrome, and Chediak–Higashi syndrome.

Histologic examination of the teeth revealed normal tooth structure (Fig 2). Hematologic examination showed periodic alteration of alkaline phosphatase and normal blood
values. Plaque test results showed a pattern of colonization relevant to the patient’s age. *Aggregatibacter actinomycetemcomitans* was not detected. These findings were inconclusive for a final diagnosis of the conditions and syndromes mentioned earlier.

At the scheduled appointment for dental treatment under GA, periodontal examination revealed mild gingival inflammation and plaque accumulation but no attachment loss. Permanent rehabilitation of the patient was performed (Fig 3), followed by tooth cleaning and fluoride treatment. As shown in Fig 3, all teeth had normal enamel structure. A panoramic radiograph was acquired to evaluate bone structure (Fig 4). Through clinical and radiographic evaluation, vertical bone loss and accelerated dental development were observed. More specifically, the first permanent lower right molar, which normally erupts at 6 years old, and the first upper left premolar, which normally erupts at 11 years old, were erupting at the age of 4.5 years.

At 1-year follow-up, the permanent teeth that had erupted showed class 1 mobility. The anterior teeth were vital, and no inflammation was observed in the gingival tissues. Radiographic examination revealed excessive bone destruction in the anterior region (Fig 5), and eruption of permanent upper and lower central incisors, first molars, and upper premolars at the age of 5.5 years was also recorded (Fig 6). A blood test for bone metabolism markers revealed an elevated RANKL serum level (1459 pg/mL vs RANKL normal values of 983–1207 pg/mL) and a lower-than-normal OPG serum level (3213 pg/mL vs OPG normal values of 3996–5808 pg/mL). Vitamin D deficiency was also apparent. The patient’s genome was checked for RANK mutations, but no findings were reported.

After consultation with the endocrinologist, the child was treated with 25(OH) vitamin D3 supplements daily (30 ng/mL) for 6 months, and the tooth mobility has been absent ever since. The patient is monitored by the endocrinologist to avoid future bone destruction.

**DISCUSSION**

This report suggests that early exfoliation of primary teeth can be a side effect of bone resorption in patients with CAH. Early exfoliation of the primary dentition and bone loss is an uncommon finding in children. However, whenever it occurs is important to diagnose the patient early because it is usually correlated with serious systematic diseases.¹⁸
Premature loss of primary teeth has not been reported in the past in a case of CAH. However, CAH should be included in the differential diagnosis of premature loss of teeth because disturbance in dental development may be the first symptom of this disorder. Other systematic diseases that may cause premature exfoliation of teeth include hypophosphatasia, aggressive periodontitis, Papillon–Lefèvre syndrome, Singleton–Merten syndrome, Hajdu–Cheney syndrome immuno-deficiency (neutropenia, leukocyte adhesion deficiency, leukemia), Langerhans histiocytosis, diabetes mellitus, hyperthyroidism, cherubism, dentinal dysplasia, Ehlers–Danlos syndrome, Coffin–Lowry syndrome, Chediak–Higashi syndrome, acatalasia, acrodynia, mucocutaneous dyskeratosis, and erythromelalgia. Hypophosphatasia, immunodeficiency, diabetes mellitus, hyperthyroidism, and acatalasia can be excluded with hematologic examination. In the present case, blood tests revealed normal blood values in all tested enzymes, hormones, or cell numbers. Langerhans histiocytosis, cherubism, and dentinal dysplasia present with bone lesions or dental morphology characteristic for these type of diseases. Acrodynia is usually correlated with mercury poisoning, which was not recorded in the patient's medical history. Papillon–Lefèvre syndrome, Ehlers–Danlos syndrome, mucocutaneous dyskeratosis, and erythromelalgia present specific dermatologic findings that were absent in the present patient.

Children with Coffin–Lowry syndrome have mental retardation, distinguishing facial characteristics, and skeletal abnormalities that were not present in this child. Moreover, none of the clinical features of Chediak–Higashi syndrome, such as albinism, strabismus, and nystagmus, infections of the skin and respiratory tract, and characteristic alterations in blood cell values and cell morphology, were found in the present patient. Patients with Singleton–Merten syndrome usually present with skeletal and joint defects, glaucoma, and calcification of the aortic arch, whereas Hajdu–Cheney syndrome is characterized by joint problems and multiple osteolytic lesions. None of these defects were apparent in this patient.

Aggressive periodontitis is a rare condition characterized by rapid loss of periodontal attachment and destruction of the bone. Usually the patients have an unrevealing medical history and present with minor inflammation of gingival tissues and minimal plaque accumulation that is incompatible with the severity of periodontal damage. Sometimes high levels of A. actinomycetemcomitans are detected. In the present patient, A. actinomycetemcomitans was not detected in the plaque microflora. The patient presented with bone destruction and gingival inflammation, but loss of attachment was not observed. However, attachment loss measurement cannot be clinically reliable in children because of continuous growth and tooth eruption.

The present case reported a child with severe bruxism. Current literature suggests a prevalence of 5.9% to 49.6% of bruxism in children. However, tooth loss due to bruxism has not been reported in the literature in either children or adults despite the high prevalence of bruxism. Bone destruction has been correlated with adrenal gland hormone.
alterations. Usually children with CAH present with accelerated bone growth caused by androgen overproduction. However, osteoclastogenesis has also been reported in these children as a side effect of long-term glucocorticoid therapy. This might have caused the bone destruction in this patient. In addition, osteoclastogenetic activity can be monitored by testing the RANKL and OPG levels in the patient’s serum. In patients with CAH, RANKL levels are elevated whereas OPG levels are lower than normal. The findings in the present case report are consistent with the literature and suggest that patient had elevated osteoclastic activity. For this reason, the endocrinologist suggested a serum test for vitamin D, which was found to be lower than normal, and the patient was given supplements to increase bone density. Other conditions presenting with bone destruction have been reported in the literature. In particular, familial expansive osteolysis and Paget disease of the bone have been related to mutations in the gene responsible for receptor activator of nuclear factor-κB (RANK). However, in the present case no RANK mutations were detected in the genome. Moreover, RANKL levels have been correlated with disturbances in tooth eruption. Accelerated development has been reported in children with CAH, as was observed for this patient. Accelerated eruption of permanent teeth may have been caused by corticosteroid- or androgen-induced bone destruction and can be a sign of bone metabolism disturbances. However, premature exfoliation in the present case may have been a side effect of accelerated eruption of permanent teeth. Additional case-control studies will be needed to define the correlation of premature exfoliation and bone resorption in patients with CAH.

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