Clinical and radiographic features of Hutchinson-Gilford progeria syndrome: A case report

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Abstract

Hutchinson-Gilford progeria syndrome (HGPS) is a rare dysmorphic syndrome characterized by several features of premature aging with clinical involvement of the skin, bones, and cardiovascular system. HGPS has an estimated incidence of one in four million to one in eight million births. The main clinical features of HGPS include short stature, craniofacial dimorphism, alopecia, bone fragility, and cardiovascular disorders. The most frequent cause of death is myocardial infarction at a mean age of 13 years old. Dental manifestations include delayed development and eruption of teeth, discoloration, crowding and rotation of teeth, and displaced teeth. Cone beam computed tomography images revealed the absence of the sphenoid, frontal, and maxillary sinus, flattening of the condyles and gelenoid fossa, and bilateral hypoplasia of the mandibular condyles. The disease is caused by mutations in lamin A/C (LMNA). Here, we present a case report of an 11-year-old boy with classical features of HGPS, which was caused by a de novo germ-line mutation (C1824T, G608G) in exon 11 of the LMNA gene. Some uncommon HGPS-associated features in our patient, such as alterations in the facial sinuses and hypoplasia of the condyles, contributed to the expansion of the phenotypic spectrum of this syndrome from a dentomaxillofacial perspective.
INTRODUCTION

Hutchinson-Gilford progeria syndrome (HGPS; OMIN #176670) is an uncommon genetic disorder characterized by accelerated aging with clinical involvement of the skin, bones, and cardiovascular system[1-3]. The prevalence of HGPS is one in four million to one in eight million live births; males are more frequently affected than females, and the intellect of the affected children is unimpaired[4]. Clinically, individuals with HGPS demonstrate short stature, prominent eyes, micrognathia, craniofacial disproportion, loss of subcutaneous fat, alopecia, beaked nose, coxa valga, pathologic bone fractures, radiolucent terminal phalanges, hearing loss, photophobia, hypertension, hyperlipidemia, atherosclerosis, and cardiovascular disorders. The most frequent cause of death is myocardial infarction at a mean age of 13 years old[4,14]. Oral alterations include high rates of tooth decay, crowding, delayed tooth development and eruption, tooth discoloration, hypodontia, maxillary and mandibular hypoplasia, and small mouth opening[9,17-19]. Recognition of dentomaxillofacial features of HGPS may allow oral health problems to be readily identified and aid in implementation of preventative treatment plans to improve quality of life[16,19]. Although HGPS demonstrates both autosomal dominant and autosomal recessive modes of inheritance, most cases are due to sporadic mutations[16]. Mutations in the lamin A/C (LMNA) gene are responsible for HGPS[17-19].

Here we report a case of HGPS in an 11-year-old boy with an uncommon phenotype and a de novo heterozygous silent mutation at amino acid 608 (G608G) of the LMNA gene.

CASE REPORT

An 11-year-old boy with a clinical diagnosis of HGPS was referred to the Clinical Department, School of Dentistry, Federal University of Pará, Brazil for oral health care. He was suffering from angina, peptic ulcer disease, and limited joint mobility. His current medications were pravastatin (5 mg/d) to prevent cardiovascular disease and ranitidine (150 mg/d) for the treatment of the peptic ulcer. The patient had normal neurodevelopment and showed the classical clinical features of HGPS, including short stature, low weight/height ratio, thin and inelastic skin, eyes slightly open when sleeping, photophobia, osteoporosis in the femur region, generalized alopecia, prominent scalp veins, small face with a beaked nose, and high-pitched voice (Figure 1). The patient had no apparent hearing loss. Echocardiogram and electrocardiogram results, blood pressure, pulse, and oxygen saturation were within normal limits. A hand-wrist radiograph showed radiolucencies of the terminal phalanges and the skeletal maturity of a 14-year-old boy with a chronological age of 11 years and 8 mo.

His height and weight were 1.1 m and 17.4 kg, respectively, which is well below the 3rd percentile for his age and only 4.4 kg greater than expected for a normal 4.5-year-old boy. Oral examination revealed micrognathia, class II malocclusion, and chronic trimus. Erupted teeth were of normal size, shape and color, but the permanent incisors were lingually erupted. The patient had gingivitis and low salivary flow, but had no dental caries and brushed his teeth while supervised by the mother. An orthopantomographic radiograph showed reduced dimensions of both arches with consequent lack of space for the correct positioning of the permanent teeth, mandible with a steep mandibular angle, eruption of the permanent teeth, and congenitally missing left upper second premolar and both lower second premolars (Figure 2). To better visualize the craniofacial features, cone beam computed tomography (CBCT) was performed. CBCT images revealed the absence of the sphenoid, frontal, and maxillary sinuses (Figure 3A and B), flattening of the condyles and glenoid fossa, and bilateral hypoplasia of the mandibular condyles (Figure 3C). Panoramic and axial images confirmed the dental alterations (Figure 3D-F).

To confirm the clinical diagnosis of HGPS, DNA sequence analysis was performed. The parents gave informed consent before the genetic study began. Mutation analysis of the LMNA gene with genomic DNA extracted from oral mucosa cells was performed according to a published protocol[20]. The patient demonstrated a heterozygous C-to-T transition at nucleotide 1824 in exon 11 of LMNA, which created a silent point mutation at codon 608 (GGC>GGT, G608G) (Figure 4). A similar mutation was not observed in the patient’s parents or sister.

DISCUSSION

This case highlights some common and uncommon dentomaxillofacial features associated with HGPS. Tooth size was essentially normal but the eruption sequence was complicated by both incomplete mandibular and
maxillary growth and micrognathia, which contributed to dental impactions. Despite radiographic evidence of normal root development, tooth eruption appeared to be delayed by three years. In addition, the permanent incisors had erupted lingually, and two premolars were absent (hypodontia). Delayed eruption, malocclusion associated with lower anterior dentition crowding, and hypodontia are consistent findings in patients with HGPS, as well as enamel hypoplasia and discoloration\[^{15,20-22}\]. The permanent teeth in the current case were macroscopically normal in shape and color. Patients who do not present alterations in the joints of the hands can carry out oral hygiene perfectly, but adult supervision is required along with the use of a toothbrush with a small head due to the small oral cavity and limited mouth opening. Interestingly, CBCT images revealed the absence of the sphenoid, frontal and maxillary sinuses, shallow glenoid fossae, and bilateral hypoplasia of the mandibular condyles and articular eminences. After evaluating radiographs of 21 children aged newborn to 14.6 years old, Gordon et al\[^{14}\] concluded that articulation deformities are not a common feature of HGPS. Chen et al\[^{11}\] reported a similar case to ours and highlighted that craniofacial anomalies of HGPS contribute to increased number of caries, severe malocclusion, and problems with swallowing, feeding, and speech. However, Ullrich et al\[^{23}\] evaluated 25 patients with HGPS and identified short mandibular rami in combination with flattened mandibular condyles, shallow glenoid fossae, and hypoplastic or absent articular eminences. The significance of the sinus alterations was unclear, however, patient- and parent-related chronic trismus can occur after a long period of regular dental

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**Figure 2** Panoramic radiograph showing delayed eruption of the permanent teeth, hypodontia of the left upper second premolar and both lower second premolars, and temporomandibular joint malformation.

**Figure 3** Craniofacial features of the Hutchinson-Gilford progeria syndrome patient detected by cone beam computed tomography. A, B: On sagittal (A) and coronal (B) images, hypoplasia of the sphenoid, frontal and maxillary sinuses was evident; C: This view depicts the temporomandibular joint alteration, which was characterized by flattening of the condyle and glenoid fossa and bilateral hypoplasia of the condyles; D: Panoramic view of the cone beam computed tomography (CBCT) revealing impaction of several permanent molars and hypodontia of the premolars; E, F: Axial slices of the CBCT showed malocclusion and lingual eruption of permanent teeth.
Hutchinson-Gilford progeria syndrome (HGPS) is an uncommon genetic disorder characterized by accelerated aging with clinical involvement of the skin, bones, and cardiovascular system. Since HGPS patients develop severe atherosclerosis and death usually occurs as a result of the complications of cardiac or cerebrovascular diseases during adolescence, early diagnosis of HGPS is important. To promote survival of HGPS patients, annual analysis of the vascular status is recommended using baseline electrocardiogram, echocardiogram, and carotid duplex scans to evaluate stenosis and intimal thickness. Additional tests include a skeletal X-ray to evaluate common associated features (e.g., acroosteolysis, clavicular resorption, and coxa valga), dual-energy X-ray absorptiometry to assess bone mineral density, standard goniometry to assess global joint mobility, and nutritional assessment to optimize caloric intake.

In summary, we report one patient affected by HGPS who demonstrated unusual features, including the absence of the sphenoid, frontal and maxillary sinuses and bilateral hypoplasia of the mandibular condyles. Proper characterization of the clinical features and genetic defects is of utmost importance for correct diagnosis and timely clinical management. Furthermore, early intervention by a multidisciplinary team can increase the quality of life and survival of HGPS patients.

**Figure 4 Detection of the LMNA mutation in the Hutchinson-Gilford progeria syndrome patient.** Shown here are portions of the DNA-sequence electropherogram of the LMNA exon 11 of the affected patient, his parents and older sister. Compared to the normal sequence, the affected patient has a heterozygous C-to-T substitution at nucleotide position 1824 in the LMNA gene, which does not change the amino acid (G608G).

**Amino acid substitution:** G608G  
**Nucleotide substitution:** C1824T

**Case characteristics**
An 11-year-old boy with a diagnosis of Hutchinson-Gilford progeria syndrome (HGPS) presented with a need for oral health care.

**Clinical diagnosis**
The patient exhibited classical clinical features of HGPS, including short stature, low weight/height ratio, thin and inelastic skin, eyes slightly open when sleeping, photophobia, generalized alopecia, prominent scalp veins, small face with a beaked nose, and high-pitched voice.

**Differential diagnosis**
Differential diagnosis included neonatal progeroid syndrome (Weidemann-Rautenstrauch syndrome), acrogeria, Cockayne syndrome, Hallermann-Streiff syndrome, gerodermia osteodystplastica, Petty-Laxova-Weidemann progeroid syndrome, and Werner syndrome.

**Imaging diagnosis**
Cone beam computed tomography (CBCT) images revealed absence of the sphenoid, frontal and maxillary sinuses, flattening of the condyles and glenoid fossa, and bilateral hypoplasia of the mandibular condyles.

**Pathological diagnosis**
DNA sequence analysis and mutation analysis of the lamin A/C (LMNA) gene was performed with genomic DNA extracted from oral mucosa cells. The patient demonstrated a heterozygous C-to-T transition at nucleotide 1824 in exon 11 of LMNA, which created a silent point mutation at codon 608 (GGC>GGT, G608G).

**Treatment**
The patient received medical and dental treatment to improve his quality of life.

**Related reports**
Recognition of dentomaxillofacial features of HGPS may allow for early identification of oral health problems and for the development of preventive treatment plans to improve quality of life.

**Term explanation**
Hutchinson-Gilford progeria syndrome is an uncommon genetic disorder characterized by accelerated aging with clinical involvement of the skin, bones, and cardiovascular system.

**Experiences and lessons**
Proper characterization of the clinical features and genetic defects of HGPS is of utmost importance for correct diagnosis and initiation of timely clinical man-
agreement; early intervention by a multidisciplinary team can increase the quality of life and survival of these patients.

**Peer review**

This article reports a case of Hutchinson-Gilford progeria syndrome in an 11-year-old boy with an uncommon phenotype and a de novo heterozygous silent mutation at amino acid 608 (G608G) in the LMNA gene.

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