Yet Another Case of Typical Hallerman Streiff Syndrome Without Mutations in the \textit{GJA1} Gene

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Introduction

Hallermann-Streiff syndrome (HSS) was first time described by Hallermann in 1948 and Streiff in 1950 who distinguished it from progeria and mandibulofacial dysostosis. Disease causing pathogenic mutation in \textit{GJA1} gene has been identified in a patient with clinical features that overlap Hallermann-Streiff syndrome (HSS; MIM# 234100) with Oculodentodigital dysplasia (ODDD; MIM# 164200) [1] However none of the cases with typical HSS are found to contain mutations in the \textit{GJA1} gene. We describe an Indian patient with typical HSS first time without identifiable \textit{GJA1} mutations.

Clinical Brief

Seven year old girl of Asian- Indian origin presented to us with complaints of short stature and sparse scalp hair. She is the first child of non-consanguineous marriage. Mother’s and father’s age were 23 and 25 years respectively at the time of birth. During antenatal period mother had gestational diabetes mellitus controlled with dietary advice and medications. Proposita was born by caesarian section with birth weight of 2.5 kg (5th centile). She was operated for bilateral eye cataract and was prescribed positive pressure ventilation at nights. She developed obstructive sleep apnea from neonatal age. She studied in class appropriate for her age and is good in studies.

The proposita had proportionate short stature with height 87 cm (<5 centile) and weight 10 kgs (<5 centile). She has the following dysmorphic features (Figure 1): brachycephalic with frontal bossing (head circumference 48 cm), prominent scalp veins, sparse thin and curly scalp hair, sparse eyebrows with no eye lashes or body hair, microphthalmia with blue sclera, down slanting palpebral fissures with telecanthus, beaked nose with hypoplastic alae nasi, abnormal dentition leading to dental caries and micrognaithia. The patient fulfilled the criteria of HSS and \textit{GJA1} gene analysis was carried out.

Methods

Informed written consent was taken and blood was collected for molecular analysis. DNA was extracted with Qiaegen midi kit following manufacturer instructions. The coding regions of \textit{GJA1} gene were amplified by PCR using primers described by Pizzuti et al. [1], \textit{GJA1} gene comprises of two exons. Forward primer 5’-GATCTTTTCTTCTGTGCGG-3’ (within intron 1) and reverse primer 5’-CTCTTTCCCTAACCCTGGC-3’ yielded 925 bp product. Another set of primers 5’-TTCTCTCTGCCCCACC-3’ and 5’-GGCCTAGAAAAGCTTACCTTT-3’ (extend into the 3’ UTR) yielded 1,079-bp product. PCR conditions were standardized for both sets of primers. PCR products were sequenced in the forward and reverse directions on an ABI prism 3130 automated sequencer. Alw44I restriction enzyme analysis was done for \textit{GJA1} c.227G>A mutation.

Results

Restriction enzyme digestion failed to show the mutation reported by Pizzuti et al. [1]. Sequence analysis of the \textit{GJA1} gene showed normal sequence.

Discussion

HSS (#234100) is characterized by craniofacial deformity with eye abnormality, proportionate short stature and hypotrichosis. Our patient had all seven diagnostic features and was thus “full-blown” phenotype. Snaepen et al. [3] describes three cases of HSS and mentions it as a classification overlap with ODDD. Homozygous \textit{GJA1} gene mutation has been reported by Pizzuti et al. [1] in one of the HSS/ODDD overlap cases reported by Snaepen. This case was rather a misdiagnosis of HSS in a case of ODDD [4].

\textit{GJA1}, a gene for Cx43 is described as causative factor for ODDD [2]. \textit{GJA1} codes for connexin-43, a transmembrane protein that facilitate cell-to-cell adhesion and provide pathways for direct intercellular communication. They help in cell adhesion and migration. They are expressed in various tissues such as the heart, liver, lymphoctes, limbs etc. and play a key role in tissue homeostasis and regulation of growth, development, and differentiation. Defect in connexin availability at different period of embryogenesis might lead to developmental abnormalities of organs pertaining to the stage of development [5].

Including this study \textit{GJA1} screening has been performed in eight cases.
well-characterized patients with HSS. Paznekas et al. [2] did molecular analysis of six patients with typical HSS and no changes in the sequences of the GJA1 gene were found. None of the patients including ours diagnosed with HSS have been found to have identifiable mutation. Though a possibility of a variation in the non-coding region of the gene that affects the expression of the gene remains, it is more likely that HSS does not come under GJA1 spectrum of disease phenotype and a different gene is responsible for this disease. Typical cases of HSS should have GJA1 gene analysis to further support this. Other candidate genes should be searched for by newer sequencing technology.

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References