Association of a Novel Nonsense Mutation in KIAA1279 with Goldberg-Shprintzen Syndrome

How to Cite This Article: Salehpour Sh, Hashemi-Gorji F, Soltani Z, Ghafouri-Fard S, Miryounesi M. Association of a Novel Nonsense Mutation in KIAA1279 with Goldberg-Shprintzen Syndrome. Iran J Child Neurol. Winter 2017; 11(1):70-74.

Abstract
Goldberg-Shprintzen syndrome (OMIM 609460) (GOSHS) is an autosomal recessive multiple congenital anomaly syndrome distinguished by intellectual disability, microcephaly, and dysmorphic facial characteristics. Most affected individuals also have Hirschsprung disease and/or gyral abnormalities of the brain. This syndrome has been associated with KIAA1279 gene mutations at 10q22.1. Here we report a 16 yr old male patient referred to Center for Comprehensive Genetic Services, Tehran, Iran in 2015 with cardinal features of GOSHS in addition to refractory seizures. Whole exome sequencing in the patient revealed a novel nonsense (stop gain) homozygous mutation in KIAA1279 gene (KIAA1279: NM_015634:exon6:c.C976T:p.Q326X). Considering the wide range of phenotypic variations in GOSHS, relying on phenotypic characteristics for discrimination of GOSH from similar syndromes may lead to misdiagnosis. Consequently, molecular diagnostic tools would help in accurate diagnosis of such overlapping phenotypes.

Keywords: Goldberg-Shprintzen syndrome, KIAA1279, Mutation

Introduction
Goldberg-Shprintzen syndrome (OMIM 609460) (GOSHS) is a multiple congenital anomaly syndrome inherited in an autosomal recessive manner and first described by Goldberg and Shprintzen in 1981 (1) and referred as Goldberg-Shprintzen syndrome (2). It is distinguished by intellectual disability, microcephaly, and dysmorphic facial characteristics. The majority of affected persons suffer from Hirschsprung disease and/or gyral defects in the brain, as the results of shortcomings in migration of neurons during embryogenesis. Additionally, megalocornea or urogenital anomalies may be detected in patients. The phenotypic features resembles Mowat-Wilson syndrome (MOWS; 235730) but these two disorders are genetically distinctive (3). GOSHS has been associated with KIAA1279 gene mutations at 10q22.1 (4). KIAA1279 encodes KIF binding protein (KBP). KBP has two tetratricopeptide repeats (TPRs) that are structural motifs involved in protein–protein interactions (5). KBP has been shown to be universally and pervasively expressed in early development while overexpressed in the central and enteric nervous systems in later stages (6). It interacts with the actin and tubulin cytoskeleton. In addition, its expression directly influences neurite growth in a neuron-like cell line in accordance with the central and enteric neuronal developmental deficiencies observed in GOSHS patients (3).
To date, seven different KIAA1279 mutations have been described in GOSHS patients all of which being nonsense mutations (Table 1).

**Case Report**

The patient is a 16 yr old boy referred to Center for Comprehensive Genetic Services, Tehran, Iran in 2015. He was the first child of healthy first-cousin Iranian parents whose second child had congenital microcephaly and hypotonia and died at the age of 12 months without any specific diagnosis (Figure 1). The proband was delivered by a normal vaginal delivery at 40 weeks of gestation after an unremarkable pregnancy. His birth occipito-frontal head circumference (OFC) was 30 cm which was 3 standard deviations (SD) lower than the age, sex and ethnicity matched population average. Afterwards, hypotonia, delayed developmental milestones as well as seizures and mental retardation were reported.

A history of bilateral cryptorchidism, inguinal hernia and Hirschsprung disease was reported which were surgically corrected. At the age of 16, his OFC was 52.5 cm which was in normal range. In physical exam he had gait disturbance, facial dysmorphism, speech defect, excessive drooling, spasticity and dry eyes. Cytogenetic analysis of the patient showed numerically normal male karyotype. The brain magnetic resonance imaging (MRI) of the patient showed brain atrophy as well as corpus callosum hypoplasia.

Informed consents were obtained from parents before participation in the study in accordance with the protocol approved by local institutional Ethics Committee.

Blood samples were collected from patient and his parents in EDTA tubes. DNA was isolated using the standard salting out method. Whole exome sequencing was performed using Illumina’s Genome Analyzer for the patient with focus on 2752 OMIM disease genes (BGI-Clinical Laboratories, Shenzhen, China). The results were verified by Sanger sequencing using the ABI Prism3130 Genetic Analyzer (Applied Biosystems, Foster City, CA, USA). A novel nonsense (stop gain) homozygous mutation was detected in KIAA1279 gene (KIAA1279: NM_015634:exon6:c.C976T:p.Q326X) correlated with the observed phenotype. This variant has not been reported in generalist polymorphism databases (ExaC or exome variant server (EVS)), dbSNP and 1000 genome project. Combined Annotation Dependent Depletion (CADD) tool which is a tool for scoring the deleteriousness of single nucleotide variants as well as insertion/deletions variants in the human genome indicated that this variant would be deleterious with a score of 17.21. Targeted sequencing on the parents showed the expected segregation pattern (Figure 2).

**Discussion**

GOSHS is a condition first defined in patients with syndromic Hirschsprung disease and attributed to neural-crest developmental defects. Further reports of bilateral polymicrogyria in patients with KIAA1279 mutations implied that the encoded protein participates in cerebral cortex development as well (3).

In this report, we have demonstrated that a novel nonsense mutation in KIAA1279 gene is associated with GOSHS in a consanguineous family. Although Hirschsprung disease is not a consistent manifestation of this syndrome, the patient reported in the present study suffered from it. However, microcephaly as a frequent manifestation of this syndrome has been reported in the patient in infancy but current physical examination showed normal OFC in contrast with a previous report (7). As different mutations in the mentioned gene have been associated with distinct phenotypes, genotype-phenotype correlation can be established for diverse mutations leading to this syndrome. Previously reported mutations in this gene have been stop gain mutations leading to nonsense-mediated mRNA decay and loss of KBP function. The ubiquitous expression of KBP predicts that such mutations result in a wide range of phenotypic manifestation in various organs.

The observed phenotypic features in this patient including mental disturbances, seizures, and gastrointestinal as well as urogenital abnormalities are in accordance with this assumption. Another prominent feature of the patient has been his refractory seizure from his infancy, which had not been reported in GOSHS patients previously assumed as a distinguishing feature between GOSHS and MOWS (7).

**In conclusion**, the present report indicates that relying on phenotypic characteristics for discrimination of these two somehow similar syndromes may lead to
Acknowledgement
We thank the patient for his participation in this study. No financial support was utilized in this study.

Authors’ contribution
Shadab Salehpour designed the work and contributed in interpretation of clinical data. Feyzollah Hashemi-Gorji and Ziba Soltani performed the molecular analysis. Soudeh Ghafouri-Fard contributed in electronic search and wrote the manuscript. Mohammad Miryounesi supervised the study and revised the manuscript for important intellectual content. All authors agreed to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

Conflict of Interests:
The authors declare that there is no conflict of interest.

Table 1. Detected Mutations in KIAA1279 Gene

| Mutation                        | Exon number | Ethnicity       | Clinical features other than classical features                                                                 | Reference |
|---------------------------------|-------------|-----------------|---------------------------------------------------------------------------------------------------------------|-----------|
| c.G250T (E84X)                  | Exon 1      | British Pakistani | Coloboma                                                                                                      | (4)       |
| c.C268T (R90X)                  | Exon 1      | Moroccan        | No Hirschsprung disease in some cases, corneal hypoesthesia, strabismus, syndactyly, conductive hearing loss, proximal muscle weakness, megalocornea | (3, 4)    |
| EX2-3DEL (N143fsX1)             | Exons 2 and 3 | Pakistani       | Fetal polymicrogyria, microcephaly, hypoplastic corpus callosum, No Hirschsprung disease, No cardinal signs of GOSHS | (7)       |
| c.C599A (S200X)                 | Exon 3      | French, Moroccan | Ptosis, hyperopia, aortic valve incompetence, cryptorchidism, high palate, oligodontia, scoliosis, vesicoureteral reflux, multicystic renal dysplasia, femoral neck anteversion | (3)       |
| c.604_605delAG (R202IfsX2)      | Exon 3      | Iraqi           | Ventricular septal defect                                                                                            | (3)       |
| c.1116_1117insA (Y466X)         | Exon 7      | Not mentioned   | Foot anomalies including campodactyly and clinodactyly                                                              | (8)       |
| c.C976T(Q326X)                  | Exon 6      | Iranian         | Inguinal hernia, criptorchidism, seizure                                                                            | The present study |
Fig 1. Family pedigree showing the consanguinity of patients’ parents

Fig 2. Nucleotide change in KIAA1279 gene in patient (A) and his parents (B)
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