The many faces of \textit{KIF7}

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Mutations in \textit{KIF7}, the gene that encodes a component of the kinesin complex of anterograde intraflagellar transport in the cilia, have been reported to cause a range of phenotypes including hydrocephaly, acrocallosal syndrome and Joubert syndrome. In a cohort of patients with various neurogenetic phenotypes, we identified novel \textit{KIF7} mutations in two families that span the known phenotypic spectrum of \textit{KIF7}-related disorders. Surprisingly, we also identified a novel truncating \textit{KIF7} mutation in a third consanguineous family, in which the index presented with intellectual disability but no overt signs of ciliopathy, and his brain magnetic resonance imaging revealed an isolated dysgenesis of corpus callosum. This small cohort contributes novel pathogenic alleles of \textit{KIF7} and suggests that \textit{KIF7}-related phenotypes can include isolated dysgenesis of corpus callosum with intellectual disability, thus expanding the range of phenotypes that warrant sequencing of this gene.

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Ciliopathies are clinical disorders that trace their pathogenesis to the defects in the primary cilium, a microtubule-based antenna-like cellular organelle that is implicated in a wide array of cellular processes.\textsuperscript{1} This common pathoetiology underpins the remarkable phenotypic overlap between various ciliopathies that were once considered distinct clinical entities, and readily explains some of the recurrent phenotypic features such as retinal degeneration (caused by loss of the connecting cilium in the photoreceptors), polydactyly (caused by abnormal ciliary signaling response to the Sonic Hedgehog morphogen gradient) and cystic kidney disease (caused by abnormal cilary mechanical response to the urine flow).\textsuperscript{2}

Central nervous system involvement in ciliopathies is complex and ranges from the very mild or absent to the profound and lethal malformations with predilection to the posterior fossa as often observed in Meckel–Gruber syndrome.\textsuperscript{3} Acrocallosal syndrome (ACS) is a ciliopathy with a prominent central nervous system involvement usually in the form of absent corpus callosum and hydrocephalus in addition to polydactyly. \textit{KIF7} (Kinesin Family Member Protein 7) mutations have been found to cause a spectrum of phenotypes that collectively fall under the clinical definition of ACS.\textsuperscript{4} In this report, we emphasize the marked phenotypic variability of \textit{KIF7}-linked phenotypes ranging from the perinatally lethal to a previously unrecognized mild end of the spectrum that is limited to intellectual disability and absence of corpus callosum.

We report three cases with \textit{KIF7} mutation, one with a classical ACS diagnosis. Another case had brain features consistent with the diagnosis of Joubert Syndrome, and \textit{KIF7} sequencing revealed a pathogenic mutation making it among the very few reported Joubert cases to carry a mutation in this gene. The third case was very mild and the phenotype was one of intellectual disability and dysgenesis of corpus callosum dysgenesis but no other signs of ciliopathy.

Case 1 is a 9-months-old boy born to consanguineous parents. Birth weight was 3.2 kg and head circumference was unknown. He started having recurrent seizures at 4 months of age. Evaluation at 9 months showed global developmental delay, dysmorphic facial features, bilateral pre-axial lower limb polydactyly and hypotonia (Figure 1). Brain computed tomography revealed corpus callosum agenesis. Case 2 is a 2.5-years-old boy with a global developmental delay. Magnetic resonance imaging revealed a classical molar tooth sign, inferior vermis hypoplasia and subcortical changes in the posterior fossa and corpus callosum agenesis (Figure 1). Case 3 is a 4-year-old boy born to consanguineous parents. Delivery was complicated by perinatal asphyxia necessitating neonatal intensive care unit management. Evaluation at four years of age revealed facial dysmorphism with brachycephaly, frontal bossing, flat supraorbital ridges, prognathism, evverted lower lip and saddle nose with a tuft of hair surrounding a sinus. Two café au lait spots were noted. Magnetic resonance imaging-brain revealed mild dysgenesis of the corpus callosum (Figure 1).

Patients and available family members were recruited using an institutional review board approved protocol (KFSSHRC RAC# 2080006) and a written informed consent was obtained from all subjects (or legal guardians) prior to enrollment. Venous blood was collected in EDTA tubes for DNA extraction and molecular studies. In Case 1 where the phenotype was typical of ACS, the entire coding sequence of \textit{KIF7} and the flanking intronic segments were amplified by using primers designed by Primer3 and subsequently sequenced bidirectionally using standard Sanger sequencing (primers and PCR conditions are available upon request). Sequence analysis revealed a \textit{KIF7} homozygous truncating mutation (NM_198525.2:c.2272G>T, p.E758*) (Figure 2). In Cases 2 and 3,
candidate genes for sequencing were determined by using autozygosity mapping as described before. Briefly, the entire set of autozygous intervals per genome (autozygome) was determined for each index by using stretches of homozygosity ($\geq$ 2 Mb) as a surrogate after performing genomewide genotyping on the Axiom (Affymetrix, Santa Clara, CA, USA) single-nucleotide polymorphism platform and subsequent search for homozygous intervals by using AutoSNPa (dna.leeds.ac.uk/autosnpa). Genes within the autozygome were interrogated for candidacy based on the clinical features by using the Genomic Oligoarray and single-nucleotide polymorphism array evaluation tool v2.0. In Case 2, autozygome-guided candidate gene analysis linked this patient to KIF7, and gene sequencing analysis revealed a homozygous truncating mutation (NM_198525.2:c.3331C>T, p.R1111*). In Case 3, autozygome analysis revealed KIF7 as a candidate gene and sequencing confirmed the presence of a homozygous truncating mutation (NM_198525.2:c.2896_2897del, p.Ala966Profs*81) (Figure 2). Segregation with the disease was confirmed for all three mutations within the respective families.

KIF7 encodes a 1343 amino acid protein, a member of a family of 14 kinesin proteins known to be 'motor proteins', molecular proteins that hydrolyze adenosine triphosphate, converting its chemical energy into mechanical energy to aid their transport along microtubules. KIF7 is a cilia-associated protein that plays an important role in the Sonic Hedgehog signaling pathway. Specifically, KIF7 functions as a negative regulator of the Sonic Hedgehog pathway by preventing an inappropriate activation of Gli2, and as a positive regulator by preventing the processing of Gli3 into its repressor form.

KIF7 mutations have been implicated in the pathogenesis of ACS, a ciliopathy with strong central nervous system involvement in the form of intellectual disability with total or partial corpus callosum absence, in addition to craniofacial dysmorphism and polydactyl. Hydrolethalus can be considered a more severe form of ACS with a lethal brain phenotype encompassing hydrocephaly and anencephaly. Mutations in KIF7 have been identified in patients with this phenotype, in addition to HYLS1. Interestingly, KIF7 mutations have also been reported in Joubert syndrome albeit rarely.

Of the 39 cases of ACS reported so far, 22 of them have atypical features with regard to either a wider clinical spectrum or a milder phenotype. However, the very mild presentation of Case 3
that is limited to intellectual disability and dysgenesis of corpus callosum appears to be unique and raises an interesting question of the contribution of KIF7 to non-syndromic intellectual disability with corpus callosum agenesis. To address this, we sequenced KIF7 in 11 cases with corpus callosum agenesis and intellectual disability but did not identify any mutation. Thus, while this case demonstrates that KIF7 mutation can present very mildly as intellectual disability with corpus callosal dysgenesis, its contribution to this phenotype appears to be rare.

In summary, we show that KIF7 mutations can result in clinical phenotypes that span a wide spectrum and we expand that spectrum further to include intellectual disability with corpus callosal dysgenesis.

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COMPETING INTERESTS

The authors declare no conflict of interest.

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