Nystagmus: A FRMD7 Prospective

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ABSTRACT

Nystagmus is involuntary up and down, to and fro and oscillatory movement of eye. Nystagmus is caused by different inheritance pattern, which can be; AD (autosomal dominant), AR (autosomal recessive) and X-linked. This review elaborates the mutational canvas of FRMD7 gene involved in X-linked Nystagmus. In Pakistan the publication is either not available or no research findings has been reported for FRMD7 gene according to our knowledge.

Keywords
Nystagmus, X-linked, FRMD7, mutations, Pakistan.

Nystagmus; the involuntary movement of eye, acquired in infancy or later in life, that results in limited or reduced vision. It is also called “Dancing eyes” due to this involuntary eye movement1. During normal situation head rotates around an axis and distant visual images are held by rotating the eyes in the opposite direction on the axis2. The angular acceleration in the vestibule is sensed by the semicircular canal. These send massages to the nuclei, from here, then pass to the extraocular muscles to allow eye movement to fixate on one object along the head movement. When these canals are stimulated then eye condition is Nystagmus. The direction of ocular movement is related to the semicircular canal that is being stimulated3. It is revealed that 1 in 1,000-2,000 people are affected by congenital and acquired nystagmus4.

FRMD7

FRMD7 is associated with x-linked Idiopathic congenital nystagmus and reported by5 previously known as LOC90167, on Xq26-q27. The function of FRMD7 is still unclear. The FRMD7 gene comprises 12 exons and 714-amino acid5. According to HGMD professional more than 90 mutations has been reported in FRMD7.

FRMD7 Mutations

In different families from Italian, England, and German origin, Tarpey et al.,5 reported different (twenty two) mutations of FRMD7 gene in the families with congenital X-linked nystagmus. Thomas6 reported 3 novel mutations in FRMD7 gene. In another study Zhang7 reported in frame deletion on 3 base pair in first exon of the FRMD7 gene in Chinese family. Shiels8 reported mutation in the 6th exon of FRMD7 gene resulting in amino acid change, leucine to arginine on 142 position. While analyzing X-linked Turkish family with nystagmus, Kaplan9 reported mutation in FRMD7 8th exon, which resulted arginine to glycine on amino acid position 229. He10 reported a deletion mutation of 2 base pair deletion in 12th exon of FRMD7 in Chinese family. Thomas6 reported a leucine change into valine at 231 amino acid position in FRMD7 gene in X-linked nystagmus family. A splice site variant in the 11th intron of the FRMD7 gene has been reported by11.
Self reported novel mutation in FRMD7 gene with insertion of Adenine on 880 position (880insA) resulting extremely variable phenotype. In another study mutation analysis of the FRMD7 entire gene, A novel mutation (misses) is reported; A>G at nucleotide position 917 (c. A917G), which resulting the amino acid change of Arg for Gln at 305 position. The mutation was homozygous and heterozygous for female, hemizygous for male in affected individuals. Kumar compared the clinical and oculomotor characteristics of albinism and FRMD7 associated Nystagmus and showed differences in characteristics between nystagmus with albinism. Pendular waveform types and nystagmus frequency were higher in FRMD7 as compared to albinism, whereas anomalous head posture and strabismus were lower in FRMD7. A study published of a novel mutation in the 12th exon of FRMD7 gene in Chinese family having four base pair deletion at cDNA position 1486-1489 (TTTT) in another study 5 novel mutations identified in FRMD7 gene, at cDNA position 70, 689, 782, 812, and 910 with G>T AG deletion, G>A, G>T, and C>t, resulting in amino acid change of G24W, Ser232del, R260Q, C271F and R303X respectively. In another similar study AlMoallem elaborated the mutation canvas of FRMD7 and expended mutation findings by reporting five novel mutation and 4 already reported. Novel includes; a frame shift mutation cDNA position at 2036 deletion occurred, missense mutation at position 801 c changed into A, a splice-site mutation of 5 plus of c.497 G>A and a 1.29 Mb deletion.

**FRMD7 Mutation in Pakistan**

In Pakistan very limited study is performed for the Nystagmus. No sufficient study evidence has been found in Pakistan for FRMD7. A wide study is needed to elaborate the mutational canvas of the FRMD7 mutation in Pakistan. Despite high consanguinity rate in Pakistan a lot of genetic disorders and their variants are reported but no sufficient published data is available to study or analyze the FRMD7 mutation prevalence in Pakistan.

**DISCUSSION**

Idiopathic congenital nystagmus is more common inherited eye disorder associated with Nystagmus, and individual affected with this disorder often have eye visual defects. It is X-linked condition and FRMD7 gene’s mutations are responsible for Idiopathic congenital nystagmus.

Idiopathic congenital nystagmus can cause different problems in eye and in most cases severely affect the vision and it is the most common eye disorder affecting visual acuity. This condition is usually present in infancy or within 2 to 4 months of birth without other sensory complications.

The inheritance pattern of nystagmus is different in different cases; can be AD (Autosomal Dominant), AR (Autosomal Recessive) or x-linked. The x-linked nystagmus is considered to be most common.

Mutations in the FRMD7 gene is considered the major reason of x-linked nystagmus and this finding is being reported by number of research publication in recent times. In Pakistan, very limited research is done and extensive study should be taken to understand and extend the nystagmus prevalence and horizon in Pakistan.

**REFERENCES**


