Genetic variant screening and association study of NKX2-5 in congenital heart disease patients from north India

Abstract:

Background: Globally 1% of the live births are affected by some form of congenital heart anomaly. Genetics and environment both play a role in its causation but very little of these aspects are explored from the Indian subcontinent. One of the first and key transcription factors required for the formation of the heart during development is NKX2-5. Several mutations in this gene have been identified for CHDs. In this study, we screened for known and novel variants to understand their role in CHDs.

Methods: Two exons and flanking 3’ and 5’ UTR regions of NKX2-5 were sequenced in n=71 CHD cases, followed by case-control test of association and haplotype study.

Results: Only 3 known variants namely rs2277923 (c.63A> G), rs3729753 (c.606G>C), and rs703752 (c.61G>T) were identified in a total of n= 69 cases. Case-control test of association revealed no significant allelic or haplotypic association. A genotypic association was observed for rs703752 in a recessive model (χ² = 4.4702; p=0.03; Risk score=0.33), along with a trend of association for rs3729753 (χ² = 3.73; p=0.053; Risk score=1.68) and rs703752 (p=0.082).

Discussion: Although we did not identify any new mutations in the coding regions of NKX2-5 gene, our findings are important observations and incite for establishing the association between NKX2-5 variants and cardiac defects in the context of the north Indian population. There is a need to explore the role of other transcription factors, cardiac developmental pathways and establish their interaction and their role in disease biology in the Indian Subcontinent.
Figure: Diagrammatic representation of the structure of NKX2-5

Image showing different domains of NKX2-5 and reported SNPs in our study