SPECIAL COMMUNICATION
THE ROLE OF GENETICS IN THE UNDERSTANDING OF COMPLEX CONGENITAL HEART DISEASES
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One of the most dynamic organs in the human body is the heart. Cardiac development is regulated by two key factors including signaling and transcriptional pathways. Thus, during the development of the fetus, any factor that disrupts the normal functioning of these factors may lead toward congenital heart defects (CHDs). Congenital heart disease is a complex multifactorial disease that involves both environmental and genetic factors.1-2 It is defined as any defect that occurs during heart development either in the cardiac structure or its associated vessels.3 Globally, it is one of the most common reasons for infant mortality and one of the most common birth defects in paediatric patients. As a rough estimate, 8 infants in every thousand live births are born with this fatal disease.4

Every year, in Pakistan approximately 40,000 children suffered from CHDs.5 Clinically depending upon the disease severity congenital heart diseases can be divided into two major subgroups one is non-syndromic and the other is syndromic. And among all these defects cardiac septation defects are the most common accounting for approximately 50% of the cases.6 It can be further sub-grouped as isolated lesions or complex diseases in combination with other heart defects.3 The exact mechanism involved in the pathogenesis of the congenital heart remains poorly understood but the most probable mechanism is multifactorial. Recent investigations suggest the role of epigenetic factors, micro RNA and small non-coding RNAs in the development of congenital heart defects. Moreover, advancements in molecular techniques including next-generation sequencing (NGS) helps in further detecting the genetic causes of CHDs such as the novel single nucleotide polymorphisms (SNPs) and copy number variants (CNVs).7

The current approaches used for genetic diagnosis of paediatric patients suffering from CHDs include karyotype analysis, copy number variation analysis, next-generation sequencing, and whole-genome or whole-exome sequencing. The targeted NGS relies on the selected region of known gene of interest and compared to whole genome or exome sequencing it provides us deeper gene coverage with easy variant detection at a lower cost. It provides robust detection of deletions, insertion and single nucleotide polymorphisms which chromosomal microarray analysis (CMA) and karyotyping cannot detect. To date, many pathogenic variants in different genes such as CITD2, CHD7, ZFPM2, MYH6 and KMT2D have been investigated by using targeted NGS. While whole-exome or genome sequencing help in the discovery of genes involves in the pathogenesis of congenital heart defects as it gives us more resolution at a single base-pair level.

Thus accurate genetic diagnosis can be done by using the appropriate diagnostic techniques that can ultimately help in better patient counseling and clinical outcome.8 Furthermore, personalized medicines or finding mutations responsible for individual congenital heart disease patients can direct to better outcomes and approaches for each cardiac malformation phenotype. Thus, ultimately combined data of patients genotypic and phenotypic following well-designed guidelines will accelerate the translation of each SNP information into better treatment and clinical insights.9

Keywords: NGS, Single nucleotide polymorphisms, CHDs.

REFERENCES


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