



iJRASET

International Journal For Research in
Applied Science and Engineering Technology



INTERNATIONAL JOURNAL FOR RESEARCH

IN APPLIED SCIENCE & ENGINEERING TECHNOLOGY

Volume: 14 **Issue:** VIII **Month of publication:** August 2026

DOI: <https://doi.org/10.22214/ijraset.2026.84718>

www.ijraset.com

Call: ☎ 08813907089

E-mail ID: ijraset@gmail.com

Genetic-Maternal Environmental Interaction during Pregnancy as a Risk Factor for Congenital Heart Disease (CHD)

Prateeksha Anandan

Student (JSS International School), UAE

Abstract: Congenital heart disease (CHD) remains a major cause of neonatal mortality, and its underlying causes are still undefined. Increasing evidence shows that CHD results from complex interactions between genetic factors in the fetus and environmental exposures from the mother during pregnancy. This review brings together findings from experimental models, human studies, and molecular research to look at how these interactions affect heart development. Important mechanisms include reduced developmental strength, threshold effects, and changes in gene expression due to environmental factors. The review also compares findings from animal experiments with the challenges of human studies, highlighting gaps in exposure assessment. By focusing on the combined impact of genetic and maternal environmental factors, this review stresses the need for a comprehensive risk assessment and points out future directions for improving prevention strategies and the relevance of research in congenital heart disease.

I. INTRODUCTION

Congenital Heart Disease (CHD), established as the most prevalent neonatal malformation in about 1% of newborns, is a leading cause of infant morbidity and mortality^[1,2]. The etiology of CHD, despite genetic advances in pre-natal diagnosis and surgical procedures, remains unclear^[1]. While more than 100 genetic abnormalities have been found associated with CHD, only one-third of cases arise from simple genetic factors^[3]. In non-syndromic forms, there has been recognition of a multifactorial pathogenesis with interaction between inherited and non-inherited factors, including influence of gene-environment interactions^[1].

Approximately 10% of CHD cases arise from gestational exposure to teratogens, with maternal diabetes being the prevailing intrinsic environmental risk factor for CHD, identifying dysregulation of cardiac development processes^[2]. Gene-environment interplay has thus become crucial to extensive CHD research, emphasizing that neither genetic nor environmental influences act in isolation. Comprehending the complex interplay between genetic factors and maternal environmental influences is critical for improving the accuracy of risk assessment and advancing the quality of neonatal care.

Understanding genetic and maternal environmental interactions has transformed CHD research from just diagnosing the condition to focusing on prevention and prediction. This shift highlights the importance of improving maternal health before and during pregnancy^[1,3]. This paper provides a comprehensive review of current research on genetic-maternal environmental interactions in congenital heart disease (CHD), detailing the underlying biological mechanisms, recent scientific discoveries, and the broader implications for both clinical practice and future research directions.

II. REVIEW

A. Gene-Environment Interactions Reduce Embryonic Development

Gene-environment interactions (GEIs), specifically maternal environment, significantly affects the embryo's ability to buffer genetic or environmental disruptions. Experimental studies through mouse embryos heterozygous in certain cardiac genes (Tbx1 or Fgfr1/Fgfr2) show complete development under optimal conditions, but high vulnerability on exposure to environmental stressors. Experimental exposure to short-term gestational hypoxia and metabolic stress interplays with genetic variants (Nkx2-5), resulting in disruption of vital processes such as abnormal cardiac looping, arrested chamber development, and irregular outflow tract morphogenesis. Such interaction for prolonged periods causes structural defects or embryonic lethality^[4]. Following the principle of threshold effects, such interplay of mild factors due to abnormal maternal environments contribute to severe CHD outcomes.

B. Critical Role of Maternal Environmental Factors

Numerous maternal environmental exposures have been associated with CHD susceptibility, incorporating metabolic disorders (diabetes, obesity), hypoxic abnormalities through smoking, anemia, or high altitudes, alcohol and drug consumption, infections, dietary deficiencies (folate imbalance), and environmental toxins exposure.

Such factors disturb vital signaling pathways in cardiogenesis, such as regulation of cell proliferation, migration, apoptosis, and left-right patterning. Moreover, timing of exposure vastly affects severity, as early gestational mutations have fatal effects on disruption in cardiac development^[2,5]. Maternal environmental stressors therefore act as modifiers rather than independent risk factors, influencing execution of neonatal genetic programs during cardiac development.

C. Epigenetic Mechanisms as Mediators

Epigenetic mechanisms perform as a crucial link between maternal environmental exposures and alternation in cardiac gene expression. Environmental stressors induce modifications in DNA methylation, histone modifications, and non-coding RNA expression, altering transcriptional genetic programs without affecting the existing DNA sequence.

Such epigenetic alterations persist throughout the development processes, influencing cardiogenic networks and increasing CHD risks^[1]. Such mechanisms indicate how maternal exposures during gestation have long-lasting effects on fetal cardiac development. Such epigenetic changes further explains variable expression of CHD despite genetic similarities among individuals.

D. Variable Penetrance and Phenotypic Heterogeneity

The concept of genetic-maternal environment interaction effectively explains phenotypic heterogeneity in individuals having identical genetic mutations. Such individuals may display varied cardiac phenotypes, or even remain unaffected, despite genetic similarities.

Environmental exposures influence gene dosage sensitivity, indicating the effects of maternal environment on functioning of a genetic variant. Experimental models indicate that abnormal conditions may result in transformation of low-penetrance variants into disease-causing mutations^[2,4]. Such frameworks explain high frequency of sporadic CHD clinical cases.

E. Evidence from Human and Experimental Models

Human epidemiological studies along with results from animal models indicate the elevation of CHD susceptibility with interplay of adverse maternal conditions with gene expression. Population-based studies report higher prevalence of CHD among offsprings of mothers with adverse conditions during pregnancy, including metabolic disorders, hypoxia, or fatal lifestyle pollutants.

Modern genomic studies and research further highlight that interaction between maternal conditions and fetal genetic networks help accurately determine cardiac development, rather than individual factors^[2,6]. Such analysis of experimental and human data emphasizes on gene-environment interaction in pathogenesis of CHD.

F. Implications for Prevention and Risk Assessment

Recognizing the significance of genetic-maternal environmental interaction in genetic susceptibility to CHD has significant implications for preventive measures and public health. Recent advances in research has prioritized maternal health before and during pregnancy, mitigating exposure to environmental risk factors, helping identify genetically susceptible cases early.

This integrated approach enhances development of risk assessment strategies, improves prenatal consultations, and targets public health campaigns^[3,5,6]. Comprehending such interactions establishes strategic frameworks reducing CHD prevalence through early preventive measures during gestation rather than treatment.

III. DISCUSSION

This review gathers current knowledge on how fetal genetic susceptibility and maternal environmental exposures interact in congenital heart disease (CHD). Evidence from human epidemiology, experimental models, and molecular research shows that CHD is rarely caused by isolated genetic mutations. Instead, its etiology involve complex interactions between cardiac gene and maternal environment. Factors such as diabetes, hypoxia, nutritional deficiencies, and exposure to toxins can affect epigenetic regulation and influence cardiogenesis. Understanding these interactions is vital for improving risk assessment, preventive strategies, and clinical management of CHD.

IV. KEY FINDINGS

The literature consistently indicates the multifactorial nature of Congenital Heart Disease. Genetic variants, including heterozygous or hypomorphic mutations in genes related to heart development, often remain hidden unless combined with harmful maternal environmental exposures^[1,2]. Animal research shows that environmental stressors like hypoxia or metabolic issues worsen defects in heart looping, chamber formation, and outflow tract development^[3]. Epigenetic mechanisms, such as DNA methylation and modifications to histones, mediate how maternal exposures affect gene expression. This provides a link between the environment and genetic makeup^[1,2].

Human epidemiological studies support these findings, showing that maternal metabolic disorders, smoking, alcohol use, and nutritional deficiencies increase the risk of CHD, particularly when they coincide with genetic vulnerability^[2,3]. These studies emphasize the importance of threshold effects, which explain the incomplete penetrance and variability in symptoms seen in clinical CHD cases.

A. Contrasting Evidence Across Studies

While there is a general agreement on the importance of gene-environment interactions, studies differ in how they report the roles of genetics and maternal environment. Genetic research often focuses on rare, high-impact variants, especially in syndromic CHD, while epidemiological studies usually highlight maternal environmental risk factors as major causes^[2,3]. Differences may stem from the design of studies, selection of participants, and methods of assessing exposures.

Experimental animal models clearly show controlled gene-environment interactions, but applying these findings to humans is difficult due to differences in genetic background, the complexity of environmental factors, and timing of exposures^[3]. Additionally, while epigenetic changes play a crucial role, the specific modifications differ across studies, and the exact causal links between maternal exposures and CHD symptoms are still not fully understood^[1,2].

B. Limitations in Current Literature

Despite growing evidence that gene and environment interactions play a role in Congenital Heart Disease, several limitations affect the current literature. Many studies still look at single genetic variants or isolated environmental factors. This approach fails to capture how real-world risk factors work together. Also, unclear timing of maternal exposures makes it hard to identify critical developmental periods when the embryonic heart is most at risk. The limited use of multifactorial methods also complicates efforts to understand how the maternal environment affects fetal gene expression. Although animal models offer important insights into mechanisms, they do not fully reflect the complexities of human pregnancies, which include genetic diversity and various environmental factors. Lastly, most studies focus on high-income populations, which limits how broadly the findings can apply, especially to low- and middle-income areas where environmental risk factors might differ.

C. Future of Gene-Environment Interaction Research

Future research should focus on long-term studies that combine genetic, epigenetic, and environmental data to identify high-risk gene and environment combinations. Better exposure assessment, inclusion of diverse populations, and studies on mechanisms will allow for improved risk grouping and preventive actions. Ultimately, applying these findings in clinical settings could lower CHD rates and improve outcomes for newborns.

V. CONCLUSION

This review highlights the increasing awareness of Congenital Heart Disease as a condition shaped by the interaction between genetic factors and maternal environmental influences. Evidence from genetic, experimental, and epidemiological studies shows that isolated genetic or environmental factors rarely work alone in determining heart outcomes. Instead, their combination during key stages of embryonic development has the greatest impact on disease risk.

By incorporating information about environmental risk factors, gene-environment interactions, and epigenetic regulation, this paper stresses the need for a more comprehensive approach in congenital heart disease research. This approach could improve early risk assessment, direct preventive strategies focused on maternal health, and guide future investigations to find modifiable factors that contribute to the disease burden. Ongoing study of these interactions will be crucial for turning scientific knowledge into real improvements in prenatal care and long-term outcomes for individuals affected by this condition.



REFERENCES

- [1] Vecoli C, Pulignani S, Foffa I, Andreassi MG. Genetic and epigenetic mechanisms linking maternal environmental exposures and congenital heart disease. *Curr Genomics*. 2014 Oct;15(5):390–399.
- [2] Choudhury TZ. Mechanisms underlying gene-environment interactions in congenital heart disease [dissertation]. Columbus (OH): The Ohio State University; 2024. ProQuest Dissertations & Theses. No. 31693896.
- [3] Kalisch-Smith JJ, Ved N, Sparrow DB. Environmental risk factors for congenital heart disease. *Cold Spring Harb Perspect Biol*. 2020;12(3):a037234.
- [4] Moreau JLM, Kesteven S, Martin EMMA, Lau KS, Yam MX, O'Reilly VC, et al. Gene–environment interaction impacts on heart development and embryo survival. *Development*. 2019;146(7):dev172957.
- [5] Jenkins KJ, Correa A, Feinstein JA, Botto L, Britt AE, Daniels SR, et al. Noninherited risk factors and congenital cardiovascular defects: current knowledge. *Circulation*. 2007;115(23):2995–3014.
- [6] Blue GM, Kirk EP, Sholler GF, Harvey RP, Winlaw DS. Congenital heart disease: current knowledge about causes and inheritance. *Med J Aust*. 2012;197(3):155–159.



10.22214/IJRASET



45.98



IMPACT FACTOR:
7.129



IMPACT FACTOR:
7.429



INTERNATIONAL JOURNAL FOR RESEARCH

IN APPLIED SCIENCE & ENGINEERING TECHNOLOGY

Call : 08813907089  (24*7 Support on Whatsapp)