Insights into the genetic architecture underlying complex, critical congenital heart disease

Gillian M. Blue1,2, Eddie K. K. Ip3, Michael Troup3, Russell C. Dale2, Gary F. Sholler1,2, Richard P. Harvey3,4, Sally L. Dunwoodie3,4, Eleni Giannoulatou3,4, David S. Winlaw5

1 Heart Centre for Children, The Children’s Hospital at Westmead, Sydney, Australia; 2 Sydney Medical School, The University of Sydney, Sydney, Australia; 3 Victor Chang Cardiac Research Institute, Sydney, Australia; 4 St Vincent’s Clinical School, UNSW Sydney, Sydney, Australia; 5 Cincinnati Children’s Hospital Medical Center, Heart Institute, Cardiothoracic Surgery, Cincinnati, Ohio, United States of America.

Background

Congenital heart disease (CHD) has a multifactorial aetiology, raising the possibility of an underlying genetic burden, predisposing to disease but also variable expression, including variation in disease severity, and incomplete penetrance. Understanding the heritable component contributing to reduced penetrance and variable expression to prevent/ameliorate disease severity in future generations, is important and relevant to the aging CHD population.

Method

Complex CHD
Requiring neonatal surgery
(n=23)

Other CHD
Post neonatal surgery
(n=39)

Whole genome sequencing

hcCHD genes*

Exome-wide

Poisson log-linear regression
compare variant number & type

Fisher’s Exact Test
Compare number of variant carriers

* n=107 established CHD genes

Results

hcCHD gene analysis: Significantly more common variants in severe CHD compared with other CHD and significantly fewer rare and low-frequency variants.

Exome-wide analysis: Significantly less rare and low-frequency variants in severe CHD cases compared with other CHD.

Gene-based variant burden: Significantly implicated pathways among rare and low-frequency variants, included ‘Autoimmune thyroid disease’ and ‘Type 1 Diabetes Mellitus’.

Conclusion

These preliminary findings suggest that the genetic architecture of complex, critical CHD is distinct from other types of CHD due to a significant increase in common variation, specifically among hcCHD genes. Further, these findings highlight associations with regulatory genes and environmental ‘stressors’ (modifiable factors affecting the fetal-placental-maternal environment) involved in the final presentation of disease. Validation in larger cohorts, may clarify the contribution of modifying variants and their effects on variable expression, specifically relating to disease severity.

Figure description: Variant effect sizes and association with disease severity where each circle represents a contributing variant, and its size the effect of the variant on the phenotype.