What are the potential benefits to our family?
The WGS technology used in this research study can capture much more information than those currently offered in the clinical diagnostic laboratory. Therefore, we may be able to help you learn more about the heart condition in your family, including the risks to other family members and what screening or treatment may be appropriate.

What does the research study involve?
Our study involves both your child and you since in genetics it is helpful to think in the context of the family. Should you choose to take part in this study here is what is involved:

• You will meet with the CGC team for a 60-90 minute research appointment. The study will be explained in detail and you will be asked for:
  - consent for participation
  - a detailed personal and family history
  - a physical exam
  - In some cases, cardiac assessment of parents (echo, ECG)
  - sample collection (bloodwork)*

• If consented, your child and in some cases you, will undergo whole genome sequencing. The results will take 12-18 months from the time of sample collection.

• Once the WGS results are available, you will have a follow up appointment for 30-60 minutes with the CGC team +/- your SickKids cardiologist to review the results.

Possible risks
WGS may show changes in a gene that predicts a risk for a person to develop another genetic condition. This is called an additional finding. These findings are unexpected and not related to why the testing was done in the first place. If an additional finding is discovered, other family members may need to be informed and offered testing. It can therefore be upsetting and stressful to learn about these additional findings.

The genetic counsellor and/or study doctor will review all possible risks of participating in this research study with you before you are asked to provide consent.

Who can participate?
If you or your child have or are at risk for heart failure, please seek further information from your cardiologist or from our Cardiac Genome Clinic genetic counsellor, Kelsey Kalbfleisch.

For more information contact:
Kelsey Kalbfleisch
Cardiac Genome Clinic Genetic Counsellor
Ted Rogers Centre for Heart Research
Clinical and Metabolic Genetics, SickKids

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In partnership with

THE PROMISE OF A HEALTHY HEART.

Cardiac Genome Project

SickKids Centre for Genetic Medicine
What is the purpose of the research?
The Cardiac Genome Clinic (CGC) study will identify genetic changes that are associated with the risk for heart failure by using a new technology called whole genome sequencing (WGS). This will help us understand how heart failure happens and how we can diagnose it and treat it earlier. Our goals are to understand 1) the genetic basis for heart failure; 2) the challenges of implementing WGS for patients with heart diseases; 3) the best approach for improving medical care using genomic data.

What is WGS?
In the past, it was only possible to test one gene at a time. This is called single gene testing. For certain health conditions, single gene testing is still done, but it can take a long time and still may not provide an explanation. Today, one test can look for harmful variants in all of your genes at once. This test is called whole genome sequencing (WGS).

Your doctor along with the CGC genetic counsellor will report the results to you. There are four possible result outcomes:
1 – positive result
2 – negative result
3 – results of unclear significance
4 – additional findings

The table explains what each of these outcomes mean and what next steps may be suggested.

### Whole genome sequencing possible results

<table>
<thead>
<tr>
<th>Possible results</th>
<th>What this means</th>
<th>Next step(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>1. Positive</strong></td>
<td>A gene change is found that is likely causing the health problem. OR A gene change is found that is not causing your child’s health problem but may provide important information. For example, the use of certain medication may not be recommended based on this result.</td>
<td>Genetic counselling and/or talking about this result with your doctor would be important to learn more about what this means.</td>
</tr>
<tr>
<td><strong>2. Negative</strong></td>
<td>No gene changes were identified.</td>
<td>Your doctor can talk with you about this, and may or may not suggest other testing.</td>
</tr>
<tr>
<td><strong>3. Results of Unclear Significance</strong></td>
<td>A change in a gene is found, but with current scientific knowledge it cannot be determined if this change is the cause of your child’s health problems.</td>
<td>Genetic counselling and/or talking with your doctor would be important to learn more about what this means. More testing may be suggested to understand this finding.</td>
</tr>
<tr>
<td><strong>4. Additional Findings</strong></td>
<td>A change in a disease-causing gene is found that is not related to why testing was ordered in the first place. This may mean your child is at risk for developing other health problems.</td>
<td>If you decide that you want to know about additional gene changes in your child, talking with your doctor and/or genetic counsellor would be important to learn more about what this means. Additional findings might lead to recommendations for your child to see other doctors, have other tests, or avoid some medications.</td>
</tr>
</tbody>
</table>

Cells, DNA and genes
In order to understand WGS better, we need to understand how our cells, DNA and genes work. Our bodies are made up of many cells, each of which contain our DNA. DNA is like a large book that has all the instructions for how our body works. This information is structured into genes that are like the chapters in a book. There are about 25,000 genes in our cells most of which contain the instructions for making proteins. Proteins are the functional and structural parts of a cell. The type of proteins made tell the cell what kind of cell it will be, such as a heart muscle cell, a brain cell, etc., and how that cell should act. Sometimes the DNA code can vary from person to person. These genetic variants help make each of us a unique individual. Genetic variants are usually harmless. Sometimes they affect minor things like hair colour but, occasionally, they can cause or put us at risk for serious health problems.