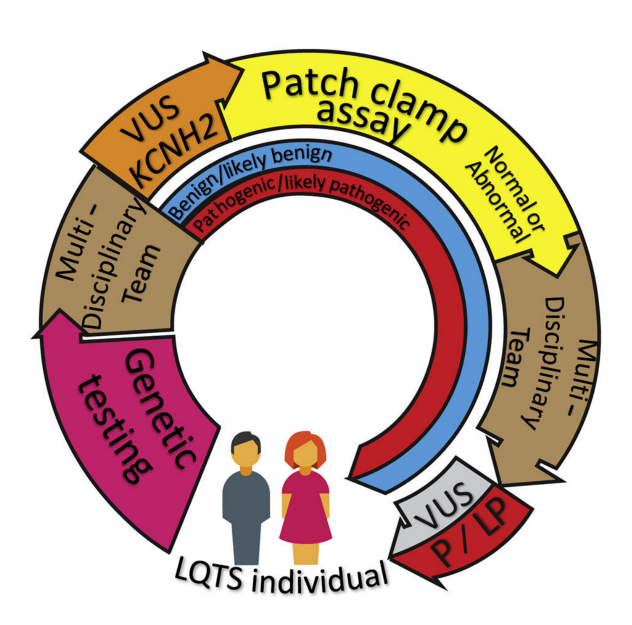


New screening test for those at risk of sudden cardiac arrest

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Graphical abstract. Credit: *The American Journal of Human Genetics* (2022). DOI: 10.1016/j.ajhg.2022.05.002

New research from the Victor Chang Cardiac Research Institute will allow families around the world to discover if they are carrying genetic mutations that cause sudden cardiac arrest—a condition that kills 9 out of 10 victims.

Researchers at the Institute have developed a new electrical test that can screen hundreds of gene mutations to pinpoint the exact mutations that are harmful to the heart for those suffering from inherited <u>heart</u> <u>disorders</u> syndromes, which can cause sudden death.

The breakthrough is a giant step forward in the accuracy and precision of genetic testing that has profound implications for not only inherited heart disorders but a wide range of neurological conditions, and muscle and kidney diseases.

Professor Jamie Vandenberg, who led the research published in two back-to-back papers in the *American Journal of Human Genetics*, says that "It's primarily <u>young people</u> with otherwise healthy hearts that die from these inherited heart disorders and even though that number is small, the consequences are long-lasting."

"When a person dies young, in the prime of their life, it's a lot more than just the death of one individual. The impact is felt on the family and their friends and that lasts forever."

Fellow author Dr. Chai-Ann Ng, of the Victor Chang Cardiac Research Institute, says that being able to identify these dangerous mutations will prevent people from dying from <u>sudden cardiac arrest</u> and ensure more



people are treated for this life-threatening disorder.

"If you can isolate the mutation and identify those at risk, there are lifestyle changes people can make, as well as taking beta-blockers or even using a defibrillator. Family members can also get themselves tested too," says Dr. Ng.

"Genetic sequencing has revealed that all of us contain a vast array of genetic variants, but we have not always been able to pinpoint if these variants are dangerous or not, only that they are different."

"So when genes are currently tested, the clinical genetics lab may tell the patient, There's a variant, but we don't know whether it raises your risk of cardiac arrest. That creates a huge amount of anxiety not just for the patient but also for the rest of the family who may also have inherited the mutation. We can now remove that uncertainty which is a big development."

Key stats

- Inherited arrhythmia disorders are found in more than half of all initially unexplained cases of sudden cardiac death in young people.
- Around 20,000 Australians suffer a cardiac arrest outside a hospital every year. Only 10% of people will survive an out-of-hospital cardiac arrest.

Professor Vandenberg's team investigated variants in genes that encode <u>ion channels</u>, which are proteins that control the movement of electrical signals between cells. The majority of genetic disorders that lead to an increased risk of sudden cardiac arrest are caused by these <u>mutations</u>.



Key findings

- In the first study, they developed a fast and accurate electrical test that assesses variants in an ion channel gene that causes an inherited heart arrhythmia condition called Long QT syndrome type 2. They're now classifying all known variants in this gene to determine which are benign and which are dangerous and will be uploading the findings to a giant genetic database that will be accessible to clinicians the world over.
- The test they have developed can easily be adapted to test other ion channel genes—not just ones associated with sudden <u>cardiac</u> <u>arrest</u> but a wide range of other diseases spanning neurological, kidney, and muscle disorders.
- In the second paper, Professor Vandenberg and his team collaborated with Dr. Kroncke at Vanderbilt University Medical Centre to develop a new method based on high throughput genome sequencing technology. This will enable them to assess the impact of every possible missense variant in KCNH2, which amounts to approximately 22,000 variants, within one to two years.

Impact

Professor Vandenberg says that they "hope that within five years, as soon as anyone gets their gene testing done, or their genomes sequenced, they will immediately find out if their <u>variant</u> is dangerous."

"It's incredible to think we will be able to screen <u>family members</u> not just across Australia but anywhere in the world and give them a diagnosis. Ultimately, this genetic database will reduce the number of cardiac arrests and deaths caused by genetic disorders."



"In the short term, it's cardiology patients who are at risk of <u>sudden</u> <u>death</u> that will benefit most. But in the longer term, the research can be adapted to assess any of the approximate 400 different ion channel <u>genes</u> in the human genome which are associated with a wide range of neural disorders, muscle and kidney problems."

More information: Jamie I. Vandenberg, A calibrated functional patch clamp assay to enhance clinical variant interpretation in KCNH2-related long QT syndrome, *The American Journal of Human Genetics* (2022). DOI: 10.1016/j.ajhg.2022.05.002

Chai-Ann Ng et al, A massively parallel assay accurately discriminates between functionally normal and abnormal variants in a hotspot domain of KCNH2, *The American Journal of Human Genetics* (2022). DOI: 10.1016/j.ajhg.2022.05.003

Provided by Victor Chang Cardiac Research Institute

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