



NON-SYNDROMIC CARDIOMYOPATHIES Requisition

Ship to:
CHEO Genetics Diagnostic Laboratory

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<https://www.cheo.on.ca/en/clinics-services-programs/genetics-diagnostic-laboratory.aspx>

ALL SECTIONS MUST BE COMPLETED

Collection Date: _____

Collection Centre: _____

CHEO Pedigree Number: _____

Patient Name: _____	_____	_____	_____
	Last	First	Initial
Health Card Number: _____			
DOB: (yy/mm/dd) _____			
Address: _____			

Telephone: _____			
Gender (circle one):	Male	Female	

Sample Requirements

Blood

Blood 2x 6 mL EDTA Blood 2x 3 mL EDTA (child) Blood 3 mL EDTA (infant ≤1 year)

For any other sample types, please contact the laboratory directly.

Health Care Provider Requesting Test

Name: _____

Registration Number: _____

Address: _____

Telephone: _____

FAX: _____

Copy to: Name: _____

Registration Number: _____

Address: _____

Telephone: _____

FAX: _____

Test Requested (see next page for the clinical testing criteria and a list of the genes included in each panel)

- ARVC panel (7 genes; see page 2 for details)
- Pan Cardiomyopathy panel (30 genes; see page 2 for details) **Note: most appropriate for DCM, or overlapping or atypical phenotypes.**
- Single gene testing (Specify Gene): _____
- Store DNA for future testing (DNA will be stored for 2 years then discarded; contact the laboratory directly if a longer storage term is required)

Note: there is different requisition for hypertrophic cardiomyopathy genetic testing; it is available at <https://www.cheo.on.ca/en/clinics-services-programs/requisitions-and-forms.aspx>

Family Variant Specific Test
(Include a copy of the family member's genetic test report. A positive control is recommended if testing was performed in a different lab)

Gene(s) _____

Variant(s) _____

Proband name: _____

Proband date of birth: _____

Relationship to proband: _____

Clinical Information

Clinical Diagnosis: HCM (Age of dx:____) DCM (Age of dx:____) ARVC (Age of dx:____) LVNC (Age of dx:____)
 Sudden cardiac arrest < 50 years old Other: _____

Cardiovascular Features:

Hypertension (treated with medication) Yes No Unknown **Ejection Fraction <40%** Yes No Unknown

Pacemaker/ implantable defibrillator Yes No Unknown **Peripartum onset** Yes No Unknown

Ventricular tachycardia Yes No Unknown

Ethnicity (be as specific as possible; this is important as the frequency of rare DNA changes can vary between ethnic backgrounds):

- Ashkenazi Jewish Black/African East Asian European First Nations French Canadian
- Hispanic Middle Eastern South Asian Other _____

Positive Family History (1st and 2nd degree relatives only) Yes (specify below) No Unknown

HCM (Age of dx:____) DCM (Age of dx:____) ARVC (Age of dx:____) LVNC (Age of dx:____)

Sudden cardiac arrest/death < 50 years old Cardiac transplant

Other: _____

NON-SYNDROMIC CARDIOMYOPATHIES TEST DETAILS

Methodology of genetic testing:

- 1) Sequencing: next-generation sequencing analysis of coding sequences of the relevant genes and 10 base pairs immediately adjacent to each exon. In addition, several deep intronic regions are analyzed for the presence of specific clinically relevant variants.
- 2) MLPA: to detect large genomic deletions and duplications, multiplex ligation-dependent probe amplification (MLPA) is performed for certain genes.

Arrhythmogenic Right Ventricular Cardiomyopathy (ARVC) Panel

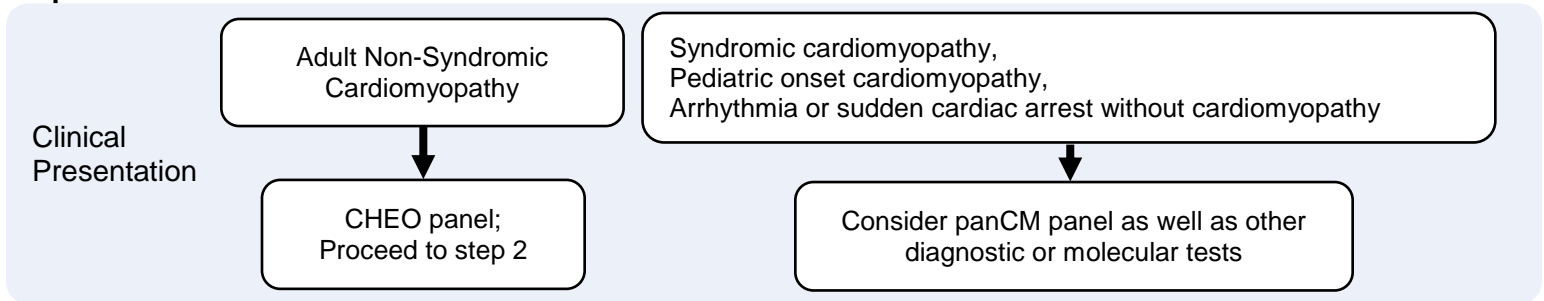
Genes included in panel: *DSC2*, *DSG2*, *DSP*, *FLNC*, *JUP*, *PKP2*, and *TMEM43* (c.1073C>T mutation only)
 Analysis includes sequencing as described above, and MLPA of *PKP2*.

Pan Cardiomyopathy Panel

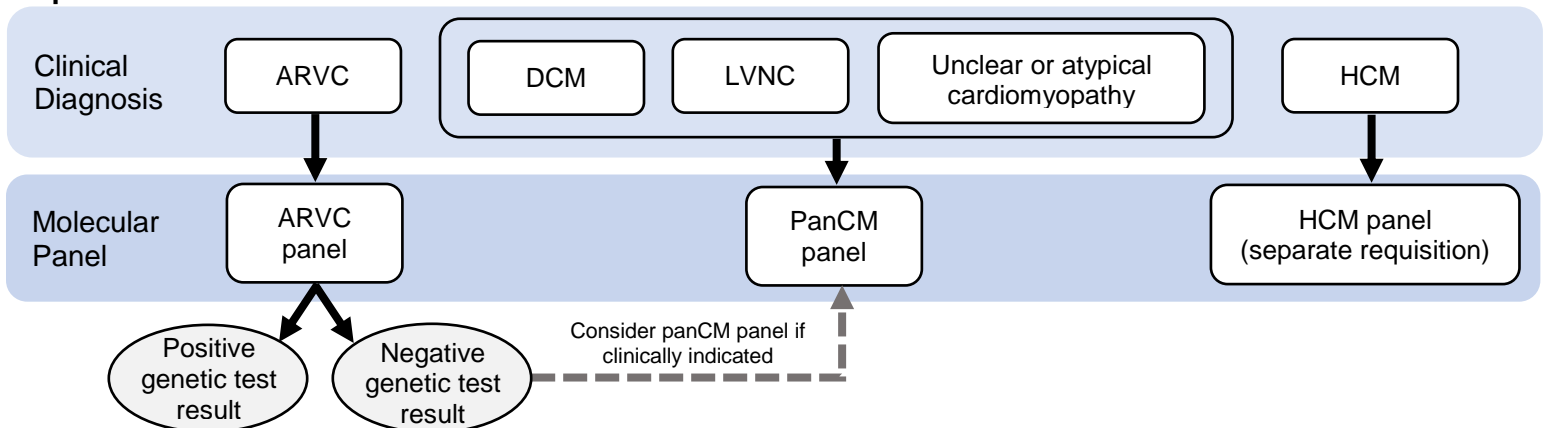
Genes included in panel: *ACTC1*, *ACTN2*, *BAG3*, *DES*, *DSC2*, *DSG2*, *DSP*, *FLNC*, *GLA*, *JUP*, *LAMP2*, *LMNA*, *MYBPC3*, *MYH7*, *MYL2*, *MYL3*, *NEXN*, *PKP2*, *PLN*, *PRKAG2*, *RBM20*, *SCN5A*, *TMEM43* (c.1073C>T mutation only), *TNNC1*, *TNNI3*, *TNNT2*, *TPM1*, *TTN*, *TTR*, and *VCL*
 Analysis includes sequencing as described above, and MLPA of *BAG3*, *MYH7*, *MYBPC3*, *PKP2*, and *TNNT2*.

Selecting a CHEO panel

Step 1:



Step 2:



Other considerations:

- **LVNC**: If additional cardiac anomalies such as congenital heart disease, consider additional tests.
- **DCM**: If extra-cardiac signs, such as muscle weakness, hearing/vision loss or if arrhythmia > cardiomyopathy, consider other etiologies and tests
- **HCM**: In patients with a family history of non-HCM cardiomyopathy or sudden cardiac death, consider the panCM panel if indicated. Consider syndromic causes of HCM, particularly in young patients with severe disease. If there is uncertainty as to which panel to order or there are additional cardiac anomalies or family history, please speak with the lab genetic counsellor.