

# Genome-wide Sequencing Ontario

## Mature Minor Secondary Findings Choices

### TYPE OF RESULT

#### PRIMARY FINDINGS

Disease-causing variants in genes that explain your medical condition

#### SECONDARY FINDINGS

Disease-associated variants in genes that are not related to the primary medical conditions for which the test has been offered, but which are associated with a high risk for other medical conditions during childhood and/or later in life.

#### What does *medically actionable* mean?

59 different genes are currently considered *medically actionable* secondary findings because if a disease-associated variant is found, there are clear medical recommendations that may be made to reduce the risk that the genetic variant will impact a person's health in the future.

Only secondary findings identified in your child can be analyzed in other family members participating in genome-wide sequencing. The choice you make regarding secondary findings will not impact the results of your child's test.

#### Actionable in Childhood

All patients under 18 will automatically have analysis of 52 genes that are medically actionable in childhood

#### All Secondary Findings

You may choose whether you wish to have analysis of all available secondary findings genes.

This includes an additional 7 genes that are not medically actionable until adulthood

For more information on Genome-wide Sequencing Ontario and other patient resources please visit [www.gsontario.ca](http://www.gsontario.ca)



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## Secondary Findings List

CONDITION	GENE	CONDITION	GENE
Adenomatous polyposis coli	APC	Loeys-Dietz syndrome	SMAD3
Aortic aneurysm, familial thoracic	ACTA2		TGFBR1
	MYH11		TGFBR2
Arrhythmogenic right ventricular cardiomyopathy	DSC2	Long QT syndrome, Brugada syndrome	KCMH2
	DSG2		KCNQ1
	DSP		SCN5A
	PKP2	Lynch syndrome	MLH1*
	TMEM43		MSH2*
	MSH6*		
Catecholaminergic polymorphic ventricular tachycardia	RYR2		PMS2*
Ehlers-Danlos syndrome, vascular type	COL3A1	Malignant hyperthermia susceptibility	CACNA1S
Familial hypercholesterolemia	APOB		RYR1
	LDLR	Marfan syndrome	FBN1
	PCSK9		
Familial hypertrophic cardiomyopathy, dilated cardiomyopathy	ACTC1	Multiple endocrine neoplasia type 1	MEN1
	GLA	Multiple endocrine neoplasia, type 2 and Familial medullary thyroid carcinoma	RET
	LMNA		
	MYBPC3	MUTYH-associated polyposis	MUTYH*
	MYH7	Neurofibromatosis, type 2	NF2
	MYL2	Ornithine carbamoyltransferase deficiency	OTC
	MYL3		
	PRKAG2	Paraganglioma-pheochromocytoma syndrome	SDHAF2
	TNNI3		SDHB
	TNNT2		SDHC
TPM1	SDHD		
Hereditary breast and ovarian cancer	BRCA1*	Peutz-Jeghers syndrome	STK11
	BRCA2*	PTEN hamartoma tumor syndrome	PTEN
Juvenile polyposis syndrome	BMPR1A	Retinoblastoma	RB1
	SMAD4	Tuberous sclerosis	TSC1
Li-Fraumeni syndrome			TSC2
		Von Hippel-Lindau syndrome	VHL
		Wilms tumor	WT1
		Wilson disease	ATP7B

\* Indicates genes that are not *medically actionable* until adulthood

