MYCODE® Results reported

Geisinger

5,545 patient-participants have received results* from the Genomic Screening and Counseling Program

For the latest results, see **geisinger.org/MyCode-results.**

May 1, 2025

350,000+	participants	have made t	the success of M	lyCode	possible

350,000+ participants have made the success of MyCode possible						
Risk Condition	Patients per condition	Z	Gene	層	Patients per gene	
				8		
CDC tie	er 1 conditions (click	link)				
Familial hypercholesterolemia (early heart attacks and strokes)	704		APOB LDLR		259 445	
Hereditary breast and ovarian cancer (early breast, ovarian, prostate, pancreatic and other cancers)	1,025	A	BRCA1 BRCA2		354 671	
Lynch syndrome (early colon, uterine and other cancers)	561		MLH1 MSH2 MSH6		52 31 238	
		ă	PMS2	A	240	
	diovascular risk	A		X		
Hereditary transthyretin amyloidosis (buildup of amyloid in the body, can lead to heart and nervous system disease)	242	8	TTR		242	
Heritable thoracic aortic disease (genetic predisposition to weakening of the wall of the aorta, leading to swelling and sometimes rupture)	51		ACTA2		51	
Inherited arrhythmias (irregular heartbeat with risk for cardiac arrest)	400		KCNH2 KCNQ1 SCN5A		44 219 137	
Inherited cardiomyopathies (diseases of the heart muscle with dangerous complications)	1,065		BAG3 DSC2 DSG2 DSP FLNC LMNA MYBPC3 MYH7 MYL2 MYL3		2 46 79 85 46 26 207 79 8 8	
			PKP2		(continued on next page)	

MyCode® results reported (continued)

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Risk Condition		Patients per condition		Gene		Patients per gene
Card	liov	ascular risk (contil	nued)		
(continued from page 1) Inherited cardiomyopathies (diseases of the heart muscle with dangerous complications)				PRKAG2 RBM20 TNNI3 TNNT2 TPM1 TTN		3 1 26 10 5 352
		Cancer risk				
Familial adenomatous polyposis (intestinal polyps and early colon cancer)		63		APC		63
Hereditary pheochromocytomas and paragangliomas (tumors that can release extra hormones and, rarely, become cancer)		114		SDHAF2 SDHB SDHC SDHD TMEM127		8 48 22 11 25
Li-Fraumeni syndrome (early breast, soft tissue, brain, adrenal and other cancers)		27	8	TP53		27
Multiple endocrine neoplasia type 1 (tumors that can release extra hormones and, rarely, become cancer)		19		MEN1		19
Multiple endocrine neoplasia type 2 (early thyroid cancer)	ğ	118	ğ	RET	ğ	118
MUTYH-associated polyposis (intestinal polyps and early colon cancer)		4		MUTYH		4
Neurofibromatosis, type 2 (noncancerous tumors in nervous system)		1		NF2		1
PALB2-related cancer risk (early onset breast, pancreatic, and ovarian cancers)		145		PALB2		145
Peutz-Jeghers syndrome (early breast, colon, pancreatic and other cancers)		2	8	STK11	8	2
Retinoblastoma (early eye cancer)		7	8	RB1		7
						(continued on next page)

MyCode® results reported (continued)

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Risk Condition		Patients per condition		Gene		Patients per gene
	Can	cer risk (continued)				
Von Hippel-Lindau syndrome (early kidney cancer and benign tumors of the brain, eye, pancreas and adrenal gland)		4		VHL		4
Wilms tumor (malignant kidney tumor)	Ď	2	A	WT1	Š	2
Misc	ella	neous phenoty	pes	S		
Biotinindase deficiency (buildup of a B vitamin in the body, can cause issues with the nervous system)		3		BTD		3
Fabry disease (enzyme defect leading to damage of blood vessels in the skin and cells in the kidneys, heart, and nervous system)		10		GLA		10
Hereditary hemochromatosis (too much iron in blood, can lead to liver and heart problems)		598		HFE		598
Hereditary hemorrhagic telangiectasia (abnormal blood vessel formation in skin, mucous membranes, lungs, liver and brain)		51		ACVRL1 ENG		13 38
Juvenile polyposis (intestinal polyps, cancer of the intestine, including colon)		3		BMPR1A		3
Juvenile polyposis / hereditary hemorrhagic telangiecstasia (intestinal polyps, cancer of the intestine, including colon/abnormal blood vessel formation in skin, mucous membranes, lungs, liver & brain)		4		SMAD4		4
Loeys-Dietz syndrome (weakening of the wall of the aorta, leading to swelling and sometimes rupture)		10		SMAD3 TGFBR1 TGFBR2		5 2 3
Malignant hyperthermia (life-threatening condition usually triggered by exposure to certain drugs used for general anesthesia)		247		RYR1		247
Marfan syndrome (connective tissue disease that can cause heart, eye, and skeletal problems)		29		FBN1		29
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Risk Condition		Patients per condition		Gene	Patients per gene
Miscella	nec	ous phenotypes	(con	tinued)	
Maturity-onset diabetes of the young (MODY) (Diabetes in the teens or early adulthood)		13		HNF1A	13
Ornithine transcarbamylase deficiency (buildup of ammonia in the blood, can cause altered mental status and seizures)		4		ОТС	4
Pompe disease (buildup of glycogen which could cause muscle probelms throughout the body)		13		GAA	13
PTEN hamartoma tumor syndrome (early breast, thyroid, uterine and other cancers, with intellectual disability in some cases)		21		PTEN	21
Retinopathy (gradual vision loss, can lead to blindness)		1		RPE65	1
Tuberous sclerosis (multiple types of benign tumors)		25		TSC1 TSC2	7 18
Vascular Ehlers-Danlos syndrome (disease of the connective tissues, including arteries and muscles, that can increase the risk for health complications, such as rupture of arteries)		14		COL3A1	14
Wilson disease (too much copper in the body, can cause liver disease and nervous system issues)		9		АТР7В	9

Totals^{†,‡}

5,617









9 9 9 5,617



 $^{^*}$ Number of patient-participants with reported results and the number per gene variant/condition $may \ not \ be \ equal \ due \ to \ the \ possibility \ of \ a \ participant \ having \ more \ than \ one \ result.$

[†]Includes some patients (~12%) already aware of their genomic result from clinical genetic testing. The process of clinical confirmation and disclosure may be modified for these patients

[‡]The gene list designated for return has shifted over time (PMID: 33576083). Totals include fewer than 10 results in genes no longer on the return list.