



Листа на 100 моногенетски болести

Болест	ГЕН	КЛАСИФИКАЦИЈА	СТЕПЕН НА СЕРИОЗНОСТ
3-Hydroxy-3-Methylglutaryl-Coenzyme A Lyase Deficiency	<i>HMGL</i>	MET	Тежок
3-Methylcrotonyl-CoA Carboxylase Deficiency 1	<i>MCCC1</i>	MET	Тежок
3-Methylcrotonyl-CoA Carboxylase Deficiency 2	<i>MCCC2</i>	MET	Тежок
Abetalipoproteinemia	<i>MTP</i>	DIG,NEUR,OPHT,HEM	Тежок
Acyl-CoA Oxidase I Deficiency	<i>ACOX1</i>	NEUR	Многу тежок
Aicardi-Goutières Syndrome	<i>SAMHD1</i>	NEUR	Тежок
Alport Syndrome, X-Linked	<i>COL4A5</i>	REN,OPHT,HEAR	Тежок
Alstrom Syndrome	<i>ALMS1</i>	OPHT,HEAR,REN,CARD	Тежок
Andermann Syndrome	<i>SLC12A6</i>	MUSC,NEUR	Тежок
Aromatase Deficiency	<i>CYP19A1</i>	SD	Умерен
Arthrogyriposis Mental Retardation Seizures	<i>SLC35A3</i>	MET	Тежок
Asparagine Synthetase Deficiency	<i>ASNS</i>	NEUR	Многу тежок
Aspartylglycosaminuria	<i>AGA</i>	MET,NEUR	Тежок
Autosomal Recessive Polycystic Kidney Disease	<i>PKHD1</i>	REN	Тежок
Bardet-Biedl Syndrome (BBS1-related)	<i>BBS1</i>	OPHT,MET,END	Тежок
Bardet Biedl Syndrome (BBS12-related)	<i>BBS12</i>	OPHT	Тежок
Beta Thalassemia	<i>HBB</i>	HEM	Многу тежок
Biotinidase Deficiency	<i>BTD</i>	MET	Тежок
Canavan Disease	<i>ASPA</i>	NEUR	Тежок
Carpenter Syndrome	<i>RAB23</i>	SKEL	Многу тежок
Choreacanthocytosis	<i>VPS13A</i>	NEUR	Многу тежок
Choroideremia, X-Linked	<i>CHM</i>	OPHT	Тежок
Citrin Deficiency	<i>SLC25A13</i>	MET	Умерен
Combined Oxidative Phosphorylation Deficiency 3	<i>TSM</i>	NEUR,MET,CARD	Многу тежок
Congenital Disorder of Glycosylation, Type 1A (PMM2-related)	<i>PMM2</i>	MET	Тежок
Congenital Neutropenia (HAX1-related)	<i>HAX1</i>	IMM	Тежок
Crigler Najjar Syndrome, Type I	<i>UGT1A1</i>	MET	Многу тежок
Cystic Fibrosis *	<i>CFTR</i>	RESP,DIG	Многу тежок
Factor XI Deficiency	<i>FXI</i>	HEM	Тежок
Familial Dysautonomia	<i>IKBKAP</i>	NEUR	Умерен
Fanconi Anemia, Type C	<i>FANCC</i>	IMM	Тежок
Fanconi Anemia, Type G	<i>FANCG</i>	HEM	Тежок
Gaucher Disease	<i>GBA</i>	NEUR,HEP,CARD	Тежок
Glutaric Acidemia, Type 2A	<i>ETFA</i>	MET	Умерен
Glycine Encephalopathy (GLDC-related)	<i>GLDC</i>	MET	Многу тежок
Glycogen Storage Disease, Type 1A	<i>G6PC</i>	MET	Умерен
Glycogen Storage Disease, Type 1B	<i>SLC37A4</i>	MET	Умерен
Glycogen Storage Disease, Type 3	<i>AGL</i>	MET	Тежок
Glycogen Storage Disease, Type 7	<i>PFKM</i>	MET	Тежок
GRACILE Syndrome	<i>BCSIL</i>	MET	Многу тежок
Hereditary Fructose Intolerance	<i>ALDOB</i>	MET	Умерен
Homocystinuria, Type cblE	<i>MTRR</i>	MET	Тежок
Hydroletharus Syndrome	<i>HYLS1</i>	NEUR,CARD	Многу тежок
Inclusion Body Myopathy, Type 2	<i>GNE</i>	MUSC	Умерен
Isovaleric Acidemia	<i>IVD</i>	MET	Тежок
Joubert Syndrome, Type 2	<i>TMEM216</i>	NEUR	Тежок
Junctional Epidermolysis Bullosa, Herlitz Type	<i>LAMC2</i>	SKIN	Тежок
Lamellar Ichthyosis, Type 1	<i>TGM1</i>	MET	Умерен
Leber Congenital Amaurosis (LCA5-related)	<i>LCA5</i>	OPHT	Тежок
Leigh Syndrome, French-Canadian Type	<i>LRPPRC</i>	NEUR,MUSC	Тежок
Leukoencephalopathy with Vanishing White Matter	<i>EIF2B5</i>	NEUR	Тежок

БОЛЕСТ	ГЕН	КЛАСИФИКАЦИЈА	СТЕПЕН НА СЕРИОЗНОСТ
Leydig Cell Hypoplasia [Luteinizing Hormone Resistance]	LHCGR	SD	Умерен
Limb Girdle Muscular Dystrophy, Type 2E	SGCB	MUSC	Тежок
Lipoamide Dehydrogenase Deficiency [Maple Syrup Urine Disease, Type 3]	DLD	MET	Тежок
Lipoprotein Lipase Deficiency	LPL	MET	Умерен
Long Chain 3-Hydroxyacyl-CoA Dehydrogenase Deficiency	HADHA	MET	Тежок
Lysinuric Protein Intolerance	SLC7A7	MET	Тежок
Maple Syrup Urine Disease, Type 1B	BCKDHB	MET	Тежок
Methylmalonic Acidemia (MMAA-related)	MMAA	MET	Многу тежок
Methylmalonic Aciduria, Type Mut(0)	MUT	MET	Тежок
Methylmalonic Aciduria and Homocystinuria, Type cbIC	MMACHC	MET	Тежок
Methylmalonic Aciduria and Homocystinuria, Type cbID	MMADHC	MET	Тежок
Mucopolysaccharidosis, Type II [Hunter Syndrome], X-Linked	IDS	RESP,CARD	Многу тежок
Mucopolysaccharidosis, Type IIIC [Sanfilippo C]	HGSNAT	MET,NEUR,OPHTH	Тежок
Multiple Sulfatase Deficiency	SUMF1	MET	Многу тежок
Myotubular Myopathy, X-Linked	MTM1	MUSC	Тежок
Navajo Neurohepatopathy [MPV17-related Hepatocerebral Mitochondrial DNA Depletion Syndrome]	MPV17	NEUR	Тежок
Neuronal Ceroid Lipofuscinosis (CLN8-related)	CLN8	NEUR	Многу тежок
Neuronal Ceroid Lipofuscinosis (MFSD8-related)	MFSD8	NEUR	Многу тежок
Neuronal Ceroid Lipofuscinosis (TPP1-related)	TPP1	NEUR	Многу тежок
Nijmegen Breakage Syndrome	NBN	NEUR	Тежок
Omenn Syndrome (RAG2-related)	RAG2	IMM	Многу тежок
Ornithine Aminotransferase Deficiency	OAT	OPHTH	Умерен
Ornithine Translocase Deficiency [Hyperornithinemia-Hyperammonemia -Homocitrullinuria (HHH) Syndrome]	SLC25A15	MET	Тежок
Pendred Syndrome	SLC26A4	HEAR,END	Умерен
Peroxisome Biogenesis Disorders Zellweger Syndrome Spectrum (PEX1-related)	PEX1	MET	Тежок
Peroxisome Biogenesis Disorders Zellweger Syndrome Spectrum (PEX2-related)	PEX2	MET	Тежок
Phenylketonurea	PAH	MET	Многу тежок
Pontocerebellar Hypoplasia, Type 1A	VRK1	NEUR, MUSC	Многу тежок
Pontocerebellar Hypoplasia, Type 2D	SEPSECS	NEUR	Многу тежок
Pontocerebellar Hypoplasia, Type 2E	VPS53	NEUR	Многу тежок
Primary Ciliary Dyskinesia (DNAH5-related)	DNAH5	RESP,INF	Умерен
Primary Ciliary Dyskinesia (DNAI1-related)	DNAI1	RESP,INF	Умерен
Primary Hyperoxaluria, Type 3	HOGA1	RESP,MET	Умерен
Pycnodysostosis	CTSK	MET	Тежок
Pyruvate Dehydrogenase Deficiency (PDHB-Related)	PDHB	NEUR,MET	Тежок
Retinal Dystrophy (RLBP1-related) [Bothnia Retinal Dystrophy]	RLBP1	OPHTH	Тежок
Retinitis Pigmentosa 25 (EYS-related)	EYS	OPHTH	Тежок
Retinitis Pigmentosa 59 (DHDDS-related)	DHDDS	OPHTH	Тежок
Sanfilippo Syndrome, Type D [Mucopolysaccharidosis IIID]	GNS	MET	Тежок
Severe Combined Immunodeficiency, Type Athabaskan	DCLRE1C	IMM	Многу тежок
Severe Combined Immunodeficiency, X-Linked	IL2RG	IMM	Многу тежок
Sickle-Cell Disease	HBB	HEM	Многу тежок
Sjögren-Larsson Syndrome	ALDH3A2	MET	Тежок
Steroid-Resistant Nephrotic Syndrome	NPHS2	REN	Тежок
Stuve-Wiedemann Syndrome	LIFR	SKEL	Тежок
Tay-Sachs Disease	HEXA	MET	Многу тежок
Usher Syndrome, Type 1F	PCDH15	HEAR	Умерен
Usher Syndrome, Type 3	CLRN1	HEAR,OPHTH	Умерен
Wolman Disease	LIPA	MET,HEP	Тежок

CARD	КАРДИОЛОШКИ	DIG	ДИГЕСТИВНИ	END	ЕНДОКРИНОЛОШКИ	HEAR	СЛУХ	HEM	ХЕМАТОЛОШКИ
HEP	ХЕПАТАЛНИ	IMM	ИМУНОЛОШКИ	INF	НЕПЛОДНОСТ	MET	МЕТАБОЛНИ	MUSC	МУСКУЛНИ
NEUR	НЕВРОЛОШКИ	OPHTH	ОФТАЛМОЛОШКИ	REN	БУБРЕЖНИ	RESP	РЕСПИРАТОРНИ	SD	СЕКСУАЛЕН РАЗВОЈ
SKEL	СКЕЛЕТНИ	SKIN	КОЖНИ						

Болезта може да се класифицира во неколку видови. Наведената класификација се заснова на најчестите симптоми поврзани со секоја состојба. Степенот на сериозноста на состојбата може да варира и зависи од специфичната мутација, знаци и симптоми.

Резултатите и можните следни чекори секогаш треба да се разгледуваат во контекст на другите клинички критериуми и треба целосно да се дискутираат со вашиот доктор. Генетско советување се препорачува кога ќе се добие резултат со висок ризик.