

INVITAE METABOLIC DISORDERS AND NEWBORN SCREENING GENE PANEL TESTS

A broad menu covering the vast majority of inherited metabolic disorders that are routinely tested as part of state newborn screening programs. Our curated panels are designed by medical genetics experts based on disorders and/or analyte results.

CLINICAL AREA: METABOLIC DISORDERS, NEWBORN SCREENING & IMMUNOLOGY

METABOLIC DISORDERS NEWBORN SCREENING CONFIRMATION

Invitae Metabolic Disorders Newborn Screening Confirmation Panel
(up to 229 genes)

Primary panel (90 genes)

ABCD1	ABCD4	ACAD8	ACADM	ACADS	ACADSB	ACADVL	ACAT1	ACSF3	AHCY	ALDH4A1	ARG1
ASL	ASS1	AUH	BCKDHA	BCKDHB	BTD	CBS	CD320	CFTR	CPS1	CPT1A	CPT2
DBT	DNAJC19	ETFA	ETFB	ETFDH	ETHE1	FAH	FTCD	G6PD	GAA	GALE	GALK1
GALT	GCDH	GCH1	GLA	GNMT	GSS	HADH	HADHA	HADHB	HCFC1	HLCS	HMGCL
HPD	HSD17B10	IDUA	IVD	LMBRD1	MAT1A	MCCC1	MCCC2	MCEE	MLYCD	MMAA	MMAB
MMACHC	MMADHC	MTR	MTRR	MUT	NAGS	OAT	OPA3	OTC	PAH	PCBD1	PC
PCCA	PCCB	PPM1K	PRODH	PTS	QDPR	SERAC1	SLC22A5	SLC25A13	SLC25A15	SLC25A20	SMPD1
SPR	SUCLA2	SUCLG1	TAT	TAZ	TMEM70						

Add-on 2,4-dienoyl-CoA reductase deficiency genes (2 genes)

DECR1 NADK2

Add-on cerebral creatine deficiency genes (3 genes)

GAMT GATM SLC6A8

Add-on congenital disorders of glycosylation genes (102 genes)

ALG1	ALG11	ALG12	ALG13	ALG14	ALG2	ALG3	ALG6	ALG8	ALG9	ATP6V0A2	B3GALNT2
B3GALT6	B3GAT3	B3GLCT	B4GALNT1	B4GALT1	B4GALT7	B4GAT1	C1GALT1C1	CHST14	CHST3	CHST6	CHSY1
COG1	COG2	COG4	COG5	COG6	COG7	COG8	DDOST	DHDDS	DOLK	DPAGT1	DPM1
DPM2	DPM3	DSE	EOGT	EXT1	EXT2	FKRP	FKTN	G6PC3	GALNT3	GFPT1	GMPPA
GMPPB	GNE	ISPD	LARGE1	LFNG	MAGT1	MAN1B1	MGAT2	MOGS	MPDU1	MPI	NGLY1
NUS1	PAPSS2	PGM1	PGM3	PIGA	PIGL	PIGM	PIGN	PIGO	PIGQ	PIGT	PIGV
PIGW	PMM2	POFUT1	POGLUT1	POMGNT1	POMGNT2	POMK	POMT1	POMT2	RFT1	RPN2	SEC23A
SEC23B	SLC26A2	SLC35A1	SLC35A2	SLC35A3	SLC35C1	SLC35D1	SRD5A3	SSR4	ST3GAL3	ST3GAL5	STT3A
STT3B	TMEM165	TMEM5	TRIP11	TUSC3	XYLT1						

Add-on generalized leukodystrophies genes (6 genes)

ARSA ASPA GALC GM2A HEXA HEXB

Add-on glucose transporter type 1 deficiency gene (1 gene)

SLC2A1

Add-on glycine encephalopathy genes (6 genes)

AMT GCSH GLDC LIAS NFU1 SLC6A9

Add-on mucopolysaccharidosis type II gene (1 gene)

IDS

Add-on Niemann-Pick type C genes (2 gene)

NPC1 NPC2

Add-on pyridoxal 5'-phosphate-dependent epilepsy gene (1 gene)

PNPO

Add-on pyridoxine-responsive epilepsy gene (1 gene)

ALDH7A1

Add-on Smith-Lemli-Opitz syndrome gene (1 gene)

DHCR7

Add-on cerebrotendinous xanthomatosis gene (1 gene)

CYP27A1

Add-on 3-hydroxy-3-methylglutaryl-CoA (HMG-CoA) lyase deficiency gene (1 gene)

HMGCS2

Add-on neuronal ceroid lipofuscinosis genes (10 genes)

ATP13A2 CLN2 (TPP1) CLN3 CLN5 CLN6 CLN8 CTSD KCTD7 MFSD8 PPT1

Add-on succinyl-CoA:3-ketoacid CoA transferase (SCOT) deficiency gene (1 gene)

OXCT1

Invitae Lysosomal Storage Disorders Newborn Screening Panel (6 genes)

Primary panel (6 genes)

GAA GALC GLA IDS IDUA SMPD1

Invitae X-Linked Adrenoleukodystrophy Newborn Screening Confirmation Test (1 gene)

Primary panel (1 gene)

ABCD1

PANELS BY ANALYTE

Invitae Low C0 Test (1 gene)

Primary panel (1 gene)

SLC22A5

Invitae Elevated C0/(C16+C18) Test (1 gene)

Primary panel (1 gene)

CPT1A

Invitae Elevated C3 Panel (9 genes)

Primary panel (9 genes)

BTD HLCS MMAA MMAB MMACHC MMADHC MUT PCCA PCCB

Invitae Elevated C3-DC Test (1 gene)

Primary panel (1 gene)

MLYCD

INVITAE METABOLIC DISORDERS AND NEWBORN SCREENING GENE PANEL TESTS (continued)

PANELS BY ANALYTE (continued)

Invitae Elevated C4 Panel (3 genes)	Primary panel (3 genes) ACAD8 ACADS ETHE1
Invitae Elevated C4-OH Panel (2 genes)	Primary panel (2 genes) HADH HIBCH
Invitae Elevated C4 & C5 Panel (7 genes)	Primary panel (7 genes) ETFA ETFB ETFDH ETHE1 SLC52A1 SLC52A2 SLC52A3
Invitae Elevated C5 Panel (2 genes)	Primary panel (2 genes) ACADSB IVD
Invitae Elevated C5-DC Test (1 gene)	Primary panel (1 gene) GCDH
Invitae Elevated C5-OH Panel (12 genes)	Primary panel (12 genes) ACAT1 AUH BTD DNAJC19 HLCS HMGCL HSD17B10 MCCC1 MCCC2 OPA3 SERAC1 TAZ
Invitae Elevated C6, C8 & C10 Test (1 gene)	Primary panel (1 gene) ACADM
Invitae Elevated C14 & C14:1 Test (1 gene)	Primary panel (1 gene) ACADVL
Invitae Elevated C16-OH, C16:1-OH, C18-OH & C18:1-OH Panel (2 genes)	Primary panel (2 genes) HADHA HADHB
Invitae Elevated C16, C16:1, C18, & C18:1 Panel (2 genes)	Primary panel (2 genes) CPT2 SLC25A20
Invitae Elevated Arginine Test (1 gene)	Primary panel (1 gene) ARG1
Invitae Elevated Citrulline Panel (up to 5 genes)	Primary panel (4 genes) ASL ASS1 PC SLC25A13 Add-on dihydroipoamide dehydrogenase deficiency gene (1 gene) DLD
Invitae Low Citrulline Panel (3 genes)	Primary panel (3 genes) CPS1 NAGS OTC
Invitae Elevated Glycine Panel (up to 62 genes)	Primary panel (6 genes) AMT GLDC GCSH LIAS NFU1 SLC6A9 Add-on organic acidemia genes (56 genes) ACAD8 ACADSB ACAT1 ACSF3 ASPA AUH BCKDHA BCKDHB BTD D2HGDH DBT DHTKD1 DLD DNAJC19 ETFA ETFB ETFDH ETHE1 FBP1 FH FTCD GCDH GSS HIBCH HLCS HMGCL HSD17B10 IDH2 IVD L2HGDH MCCC1 MCCC2 MCEE MLYCD MMAA MMAB MMACHC MMADHC MUT NFU1 OGDH OPA3 OPLAH OXCT1 PCCA PCCB POLG PPM1K SERAC1 SLC13A5 SLC25A1 SLC25A19 SUCLA2 SUCLG1 TAZ TMEM70
Invitae Elevated Leucine Panel (5 genes)	Primary panel (5 genes) BCKDHA BCKDHB DBT DLD PPM1K
Invitae Elevated Methionine Panel (up to 6 genes)	Primary panel (4 genes) AHCY CBS GNMT MAT1A Add-on additional causes of elevated methionine genes (2 genes) FAH SLC25A13
Invitae Elevated Phenylalanine Panel (6 genes)	Primary panel (6 genes) GCH1 PAH PCBD1 PTS QDPR SPR
Invitae Elevated Proline Panel (2 genes)	Primary panel (2 genes) ALDH4A1 PRODH
Invitae Elevated Succinylacetone Test (1 gene)	Primary panel (1 gene) FAH
Invitae Elevated Tyrosine Panel (3 genes)	Primary panel (3 genes) FAH HPD TAT
AMINOACIDOPATHIES	
Invitae Alkaptonuria Test (1 gene)	Primary panel (1 gene) HGD
Invitae Combined Methylmalonic Acidemia and Homocystinuria Panel (11 genes)	Primary panel (11 genes) ABCD4 AMN CD320 CUBN GIF HCFC1 LMBRD1 MMACHC MMADHC TCN1 TCN2
Invitae Cystinuria Panel (3 genes)	Primary panel (3 genes) PREPL SLC3A1 SLC7A9
Invitae Disorders of Serine Biosynthesis Panel (3 genes)	Primary panel (3 genes) PHGDH PSAT1 PSPH
Invitae Glycine Encephalopathy Panel (6 genes)	Primary panel (6 genes) AMT GCSH GLDC LIAS NFU1 SLC6A9

INVITAE METABOLIC DISORDERS AND NEWBORN SCREENING GENE PANEL TESTS (continued)

AMINOACIDOPATHIES (continued)

Invitae Homocystinuria Panel (up to 19 genes)	Primary panel (4 genes)										
	CBS	MTHFR	MTR	MTRR							
	Add-on combined methylmalonic acidemia and homocystinuria genes (11 genes)										
	ABCD4	AMN	CD320	CUBN	GIF	HCFC1	LMBRD1	MMACHC	MMADHC	TCN1	TCN2
	Add-on elevated methionine genes (4 genes)										
	AHCY	CBS	GNMT	MAT1A							
Invitae Hyperphenylalaninemia Panel (6 genes)	Primary panel (6 genes)										
	GCH1	PAH	PCBD1	PTS	QDPR	SPR					
Invitae Hyperprolinemia Panel (2 genes)	Primary panel (2 genes)										
	ALDH4A1	PRODH									
Invitae Maple Syrup Urine Disease Panel (up to 5 genes)	Primary panel (4 genes)										
	BCKDHA	BCKDHB	DBT	PPM1K							
	Add-on DLD gene (1 gene)										
	DLD										
Invitae Tyrosinemia Panel (3 genes)	Primary panel (3 genes)										
	FAH	HPD	TAT								

CARBOHYDRATE DISORDERS

Invitae Galactosemia Panel (3 genes)	Primary panel (3 genes)											
	GALE	GALK1	GALT									
Invitae Glucose-6-Phosphate Dehydrogenase Deficiency Test (1 gene)	Primary panel (1 gene)											
	G6PD											
Invitae Glucose Transporter Type 1 Deficiency Syndrome Test (1 gene)	Primary panel (1 gene)											
	SLC2A1											
Invitae Comprehensive Glycogen Storage Disease Panel (23 genes)	Primary panel (23 genes)											
	AGL	ALDOA	ENO3	FBP1	G6PC	GAA	GBE1	GYG1	GYS1	GYS2	LAMP2	LDHA
	PFKM	PGAM2	PHKA1	PHKA2	PHKB	PHKG2	PYGL	PYGM	RBCK1	SLC2A2	SLC37A4	
Invitae Liver Glycogen Storage Disease Panel (11 genes)	Primary panel (11 genes)											
	AGL	FBP1	G6PC	GBE1	GYS2	PHKA2	PHKB	PHKG2	PYGL	SLC2A2	SLC37A4	
Invitae Muscle Glycogen Storage Disease Panel (up to 15 genes)	Primary panel (14 genes)											
	ALDOA	ENO3	GAA	GBE1	GYG1	GYS1	LAMP2	LDHA	PFKM	PGAM2	PHKA1	PHKB
	PYGM	RBCK1										
	Add-on PGM1 gene (1 gene)											
	PGM1											
Invitae Hereditary Fructose Intolerance Test (1 gene)	Primary panel (1 gene)											
	ALDOB											
Invitae Rare Carbohydrate Disorders Panel (2 genes)	Primary panel (2 genes)											
	FBP1	SLC5A1										

CEREBROTENDINOUS XANTHOMATOSIS

Invitae Cerebrotendinous Xanthomatosis Test (up to 3 genes)	Primary panel (1 gene)										
	CYP27A1										
	Add-on sitosterolemia genes (2 genes)										
	ABCG5	ABCG8									

CONGENITAL DISORDERS OF GLYCOSYLATION

Invitae Congenital Disorders of Glycosylation Panel (up to 103 genes)	Primary panel (49 genes)											
	ALG1	ALG11	ALG12	ALG13	ALG2	ALG3	ALG6	ALG8	ALG9	ATP6V0A2	B3GLCT	CHST14
	COG1	COG2	COG4	COG5	COG6	COG7	COG8	DHDDS	DOLK	DPAGT1	DPM1	DPM2
	DPM3	G6PC3	GFPT1	GMPPA	GMPPB	MAGT1	MAN1B1	MGAT2	MOGS	MPDU1	MPI	NGLY1
	PGM1	PGM3	PMM2	RFT1	SEC23B	SLC35A1	SLC35A2	SLC35C1	SRD5A3	SSR4	TMEM165	TRIP11
	TUSC3											
	Add-on preliminary evidence genes (11 genes)											
	ALG14	B4GALT1	DDOST	NUS1	PIGM	RPN2	SEC23A	SLC35A3	ST3GAL3	STT3A	STT3B	
	Add-on disorders of O-mannosylation genes (13 genes)											
	B3GALNT2	B4GAT1	FKRP	FKTN	GNE	ISPD	LARGE1	POMGNT1	POMGNT2	POMK	POMT1	POMT2
	TMEM5											
	Add-on glycosylation genes not involved in N-glycosylation genes (30 genes)											
	B3GALT6	B3GAT3	B4GALNT1	B4GALT7	CIGALT1C1	CHST3	CHST6	CHSY1	DSE	EOGT	EXT1	EXT2
	GALNT3	LFNG	PAPSS2	PIGA	PIGL	PIGM	PIGN	PIGO	PIGQ	PIGT	PIGV	PIGW
	POFUT1	POGLUT1	SLC26A2	SLC35D1	ST3GAL5	XYLT1						

CREATINE BIOSYNTHESIS AND TRANSPORT DISORDERS

Invitae Cerebral Creatine Deficiency Panel (3 genes)	Primary panel (3 genes)											
	GAMT	GATM	SLC6A8									

INVITAE METABOLIC DISORDERS AND NEWBORN SCREENING GENE PANEL TESTS (continued)

CYSTIC FIBROSIS

Invitae Cystic Fibrosis Newborn Screening Confirmation Test (1 gene) **Primary panel (1 gene)**
CFTR

FATTY ACID OXIDATION DEFECTS

Invitae Fatty Acid Oxidation Defects Panel (up to 22 genes) **Primary panel (18 genes)**
ACADM ACADS ACADSB ACADVL CPT1A CPT2 ETFA ETFB ETFDH HADH HADHA HADHB
HMGCL HMGCS2 MLYCD NADK2 SLC22A5 SLC25A20

Add-on preliminary-evidence gene (1 gene)
DECRI

Add-on riboflavin transporter deficiency genes (3 genes)
SLC52A1 SLC52A2 SLC52A3

Invitae Ketogenesis Disorders Panel (2 genes) **Primary panel (2 genes)**
HMGCL HMGCS2

Invitae Ketolysis Disorders Panel (2 genes) **Primary panel (2 genes)**
ACAT1 OXCT1

Invitae Medium Chain Acyl-CoA Dehydrogenase Deficiency Test (1 gene) **Primary panel (1 gene)**
ACADM

Invitae Multiple Acyl-CoA Dehydrogenase Deficiency Panel (up to 6 genes) **Primary panel (3 genes)**
ETF A ETFB ETFDH
Add-on riboflavin transporter deficiency genes (3 genes)
SLC52A1 SLC52A2 SLC52A3

Invitae Very Long Chain Acyl-CoA Dehydrogenase Deficiency Test (1 gene) **Primary panel (1 gene)**
ACADVL

LYSOSOMAL STORAGE DISORDERS

Invitae Comprehensive Lysosomal Storage Disorders Panel (up to 53 genes) **Primary panel (52 genes)**
AGA ARSA ARSB ASAH1 ATP13A2 CLN2 (TPPI) CLN3 CLN5 CLN6 CLN8 CTNS CTSA
CTSD CTSF CTSK DNAJC5 FUCA1 GAA GALC GALNS GLA GLB1 GM2A GNPTAB
GNPTG GNS GRN GUSB HEXA HEXB HGSNAT HYAL1 IDS IDUA KCTD7 LAMP2
LIPA MAN2B1 MANBA MCOLN1 MFSD8 NAGA NAGLU NEU1 NPC1 NPC2 PPT1 PSAP
SGSH SLC17A5 SMPD1 SUMF1

Add-on chitotriosidase deficiency gene (1 gene)
CHIT1

Invitae Cystinosis Test (1 gene) **Primary panel (1 gene)**
CTNS

Invitae Farber Lipogranulomatosis Test (1 gene) **Primary panel (1 gene)**
ASAH1

Invitae Fabry Disease Test (1 gene) **Primary panel (1 gene)**
GLA

Invitae GM2 Gangliosidosis Panel (3 genes) **Primary panel (3 genes)**
GM2A HEXA HEXB

Invitae Krabbe Disease Panel (up to 2 genes) **Primary panel (1 gene)**
GALC
Add-on prosaposin deficiency gene (1 gene)
PSAP

Invitae Lysosomal Acid Lipase Deficiency Test (1 gene) **Primary panel (1 gene)**
LIPA

Invitae Metachromatic Leukodystrophy Panel (up to 7 genes) **Primary panel (3 genes)**
ARSA PSAP SUMF1
Add-on generalized leukodystrophies genes (4 genes)
ASPA GALC HEXA HEXB

Invitae Mucopolipidosis Panel (4 genes) **Primary panel (4 genes)**
GNPTAB GNPTG MCOLN1 NEU1

Invitae Comprehensive Mucopolysaccharidoses (MPS) Panel (up to 23 genes) **Primary panel (11 genes)**
ARSB GALNS GLB1 GNS GUSB HGSNAT HYAL1 IDS IDUA NAGLU SGSH
Add-on mucopolipidosis and oligosaccharidoses genes (12 genes)
AGA CTSA CTSK FUCA1 GNPTAB GNPTG MAN2B1 MANBA MCOLN1 NAGA NEU1 SLC17A5

Invitae Mucopolysaccharidosis Type I (MPS I) Test (up to 6 genes) **Primary panel (1 gene)**
IDUA
Add-on clinically overlapping lysosomal storage disorder genes (5 genes)
ARSB GNPTAB GUSB IDS SUMF1

Invitae Mucopolysaccharidosis Type II Test (up to 5 genes) **Primary panel (1 gene)**
IDS
Add-on clinically overlapping genes (4 gene)
GNPTAB GUSB IDUA SUMF1

INVITAE METABOLIC DISORDERS AND NEWBORN SCREENING GENE PANEL TESTS (continued)

LYSOSOMAL STORAGE DISORDERS (continued)

Invitae Mucopolysaccharidosis Type III (MPS III) Panel (4 genes)	Primary panel (4 genes) GNS HGSNAT NAGLU SGSH
Invitae Mucopolysaccharidosis Type IV (MPS IV) Panel (up to 3 genes)	Primary panel (2 genes) GALNS GLB1 Add-on multiple sulfatase deficiency gene (1 gene) SUMF1
Invitae Multiple Sulfatase Deficiency Test (up to 16 genes)	Primary panel (1 gene) SUMF1 Add-on mucopolipidosis and mucopolysaccharidosis genes (15 genes) ARSB GALNS GLB1 GNPTAB GNPTG GNS GUSB HGSNAT HYAL1 IDS IDUA MCOLN1 NAGLU NEU1 SGSH
Invitae Comprehensive Neuronal Ceroid Lipofuscinoses Panel (up to 13 genes)	Primary panel (9 genes) CLN2(TPPI) CLN3 CLN5 CLN6 CLN8 CTSD KCTD7 MFSD8 PPT1 Add-on preliminary-evidence gene (1 gene) ATP13A2 Add-on adult-onset neuronal ceroid lipofuscinoses genes (3 gene) CTSF DNAJC5 GRN
Invitae Niemann-Pick Disease Types A and B Test (up to 2 genes)	Primary panel (1 gene) SMPD1 Add-on chitotriosidase deficiency gene (1 gene) CHIT1
Invitae Niemann-Pick Type C Panel (up to 3 genes)	Primary panel (2 genes) NPC1 NPC2 Add-on lysosomal acid lipase deficiency gene (1 gene) LIPA
Invitae Oligosaccharidoses Panel (up to 23 genes)	Primary panel (8 genes) AGA CTSA CTSK FUCA1 MAN2B1 MANBA NAGA SLC17A5 Add-on mucopolipidosis and mucopolysaccharidosis genes (15 genes) ARSB GALNS GLB1 GNPTAB GNPTG GNS GUSB HGSNAT HYAL1 IDS IDUA MCOLN1 NAGLU NEU1 SGSH
Invitae Pompe Disease Test (up to 3 genes)	Primary panel (1 gene) GAA Add-on Danon disease gene (1 gene) LAMP2 Add-on primary carnitine deficiency gene (1 gene) SLC22A5
Invitae Prosaposin Deficiency Test (1 gene)	Primary panel (1 gene) PSAP
Invitae Sandhoff Disease Test (up to 2 genes)	Primary panel (1 gene) HEXB Add-on Tay-Sachs disease gene (1 gene) HEXA
Invitae Tay-Sachs Disease Test (up to 2 genes)	Primary panel (1 gene) HEXA Add-on Sandhoff disease gene (1 gene) HEXB

METAL TRANSPORT DISORDERS

Invitae ATP7A-Related Disorders (1 gene)	Primary panel (1 gene) ATP7A
Invitae Copper Metabolism Disorders Panel (5 genes)	Primary panel (5 genes) AP1S1 ATP7A ATP7B CP SLC33A1
Invitae Wilson Disease Test (1 gene)	Primary panel (1 gene) ATP7B

NEUROTRANSMITTER DISORDERS

Invitae Neurotransmitter Disorders Panel (up to 37 genes)	Primary panel (27 genes) ABAT ALDH5A1 ALDH7A1 AMT ARHGEF9 DBH DDC GAD1 GCH1 GCSH GLDC GLRA1 GLRB GPHN MAOA PCBD1 PHGDH PNPO PSAT1 PSPH PTS QDPR SLC25A22 SLC6A3 SLC6A5 SPR TH Add-on neurodegeneration with brain iron accumulation genes (10 genes) ATP13A2 C19orf12 COASY CP DCAF17 FA2H FTL PANK2 PLA2G6 WDR45
Invitae Hereditary Hyperekplexia Panel (6 genes)	Primary panel (6 genes) ARHGEF9 CLPB GLRA1 CLRB GPHN SLC6A5

INVITAE METABOLIC DISORDERS AND NEWBORN SCREENING GENE PANEL TESTS (continued)

PURINE METABOLISM DISORDERS

Invitae Purine Metabolism Disorders Panel
(up to 10 genes)

Primary panel (9 genes)
ADA ADSL AMPD1 HPRT1 GPHN MOCOS MOCS1 PNP XDH

Add-on sulfite oxidase deficiency gene (1 gene)
SUOX

Invitae Lesch-Nyhan Syndrome Test
(1 gene)

Primary panel (1 gene)
HPRT1

PYRUVATE METABOLISM AND TRICARBOXYLIC ACID CYCLE DEFECTS

Invitae 2-Ketoglutarate Dehydrogenase Deficiency Panel (up to 4 genes)

Primary panel (3 genes)
DLD OGDH SLC25A19

Add-on alpha-ketoadipic acid dehydrogenase deficiency gene (1 gene)
DHTKD1

Invitae Citrate Transporter Deficiency Test
(1 gene)

Primary panel (1 gene)
SLC13A5

Invitae Dihydropyridine Dehydrogenase Deficiency Test (1 gene)

Primary panel (1 gene)
DLD

Invitae Fumarate Deficiency Test (1 gene)

Primary panel (1 gene)
FH

Invitae Pyruvate Carboxylase Deficiency Test
(1 gene)

Primary panel (1 gene)
PC

Invitae Pyruvate Dehydrogenase Deficiency Panel (8 genes)

Primary panel (8 genes)
DLAT DLD LIAS MPC1 PDHA1 PDHB PDHX PDP1

TREATABLE DISORDERS

Invitae Treatable Neurometabolic Disorders Panel (up to 133 genes)

Primary panel (92 genes)
ABCD1 ACAT1 AGA ALDH5A1 ALDH7A1 AMN AMT ARG1 ARSA ASL ASS1 ATP7A
ATP7B AUH BCKDHA BCKDHB BTD CBS CLN2 (TPP1) CP CPS1 CUBN CYP27A1 DBT
DHCR7 DLAT DLD ETFA ETFB ETLFDH ETHE1 GAMT GATM GCDH GCH1 GCSH
GIF GLA GLDC GLUD1 GUSB HLCS HMGCL HMGCS2 HSD17B10 IDS IDUA IVD
LIPA LMBRD1 MAN2B1 MCCC1 MCCC2 MMAA MMAB MMACHC MMADHC MOCS1 MTHFR MTR
MTRR MUT NAGS NPC1 NPC2 OTC OXCT1 PAH PANK2 PCBD1 PCCA PCCB
PDHA1 PDHB PDHX PDP1 PHGDH PNPO PPM1K PSAT1 PSPH PTS QDPR SGSH
SLC19A3 SLC25A13 SLC25A15 SLC2A1 SLC6A8 SPR TAT TH

Add-on neurometabolic conditions genes (41 genes)
ABAT ADSL AP1S1 ATP13A2 BCKDK C19orf12 CLN3 CLN5 CLN6 CLN8 COASY CTSD
D2HGDH DBH DCAF17 DDC FA2H FTL GAD1 GNS GPHN HEXA HEXB HGSNAT
HPRT1 IDH2 KCTD7 L2HGDH MAOA MFSD8 MOCOS NAGLU PLA2G6 POLG PPT1 SLC13A5
SLC33A1 SLC6A3 SUOX WDR45 XDH

Invitae Biotin-Thiamine-Responsive Basal Ganglia Disease (BTBGD) Test (1 gene)

Primary panel (1 gene)
SLC19A3

UREA CYCLE DISORDERS

Invitae Urea Cycle Disorders Panel
(up to 15 genes)

Primary panel (10 genes)
ALDH18A1 ARG1 ASL ASS1 CPS1 NAGS OAT OTC SLC25A13 SLC25A15

Add-on hyperammonemia genes (4 genes)
CA5A GLUD1 GLUL SLC7A7

Add-on hereditary orotic aciduria gene (1 gene)
UMPS

Invitae Arginase Deficiency Test (1 gene)

Primary panel (1 gene)
ARG1

Invitae Ornithine Transcarbamylase (OTC) Deficiency Test (up to 4 genes)

Primary panel (1 gene)
OTC

Add-on hereditary orotic aciduria gene (1 gene)
UMPS

Add-on low citrulline genes (2 genes)
CPS1 NAGS



INVITAE