

Uni-ECS plus Gene Panel

Disorder	Inheritance	Gene/Locus	Detection Rate	Carrier frequency	Residual risk of being a carrier	Residual risk of having an affected child *If both parents test negative	Residual risk of having an affected child *If one of the parents tests positive and one tests negative
Metabolic System							
Adrenal hyperplasia, congenital, due to 11-beta-hydroxylase deficiency	AR	<i>CYP11B1</i>	99	1/158	1/15800	1/998560000	1/63200
17-alpha-hydroxylase	AR	<i>CYP17A1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Pseudohermaphroditism, male, with gynecomastia	AR	<i>HSD17B3</i>	99	1/135	1/13500	1/729000000	1/54000
Peroxisome biogenesis disorder 1A (Zellweger)	AR	<i>PEX1</i>	99	1/500	1/50000	1/10000000000	1/200000
2,4-dienoyl-CoA reductase deficiency	AR	<i>NADK2</i>	99	<1/500	1/50000	1/10000000000	1/200000
Adrenal hyperplasia, congenital, due to 21-hydroxylase deficiency	AR	<i>CYP21A2</i>	63	1/61	1/165	1/108722	1/659
2-methylbutyrylglycinuria	AR	<i>ACADSB</i>	99	<1/500	1/50000	1/10000000000	1/200000
Adrenal hyperplasia, congenital, due to 3-beta-hydroxysteroid dehydrogenase 2 deficiency	AR	<i>HSD3B2</i>	99	<1/500	1/50000	1/10000000000	1/200000
3-Methylcrotonyl-CoA carboxylase 1 deficiency	AR	<i>MCCC1</i>	98	1/95	1/4750	1/90250000	1/19000
3-Methylcrotonyl-CoA carboxylase 2 deficiency	AR	<i>MCCC2</i>	94	1/95	1/1583	1/10027778	1/6333
3-methylglutaconic aciduria, type III	AR	<i>OPA3</i>	99	<1/500	1/50000	1/10000000000	1/200000
3-methylglutaconic aciduria, type IX	AR	<i>TIMM50</i>	89	3/500	1/1515	1/9182736	1/6061
3-methylglutaconic aciduria, type I	AR	<i>AUH</i>	99	3/500	1/16667	1/1111111111	1/66667
3-methylglutaconic aciduria, type VIII	AR	<i>HTRA2</i>	99	3/500	1/16667	1/1111111111	1/66667
3-methylglutaconic aciduria, type VII, with cataracts, neurologic involvement and neutropenia	AR	<i>CLPB</i>	99	3/500	1/16667	1/1111111111	1/66667
3-methylglutaconic aciduria, type V	AR	<i>DNAJC19</i>	99	3/500	1/16667	1/1111111111	1/66667
3-methylglutaconic aciduria with deafness, encephalopathy, and Leigh-like syndrome	AR	<i>SERAC1</i>	93	3/500	1/2381	1/22675737	1/9524
HMG-CoA synthase-2 deficiency	AR	<i>HMGCS2</i>	99	<1/250	1/25000	1/2500000000	1/100000
3-hydroxyacyl-CoA dehydrogenase deficiency	AR	<i>HADH</i>	92	1/240	1/6250	1/156250000	1/25000
Peroxisome biogenesis disorder 4A (Zellweger)	AR	<i>PEX6</i>	99	1/280	1/28000	1/3136000000	1/112000
Glycine encephalopathy	AR	<i>AMT</i>	99	1/373	1/37300	1/5565160000	1/149200
Hyperphenylalaninemia, BH4-deficient, A	AR	<i>PTS</i>	98	<1/500	1/25000	1/2500000000	1/100000
Hyperphenylalaninemia, BH4-deficient, B	AR	<i>GCH1</i>	83	<1/500	1/2941	1/34602076	1/11765
Hyperphenylalaninemia, BH4-deficient, C	AR	<i>QDPR</i>	99	<1/500	1/50000	1/10000000000	1/200000
Hyperphenylalaninemia, BH4-deficient, D	AR	<i>PCBD1</i>	99	1/200	1/20000	1/1600000000	1/80000
D-bifunctional protein deficiency	AR	<i>HSD17B4</i>	93	1/158	1/2257	1/20378776	1/9029
HMG-CoA lyase deficiency	AR	<i>HMGCL</i>	99	<1/500	1/50000	1/10000000000	1/200000
Glycogen storage disease IIIa	AR	<i>AGL</i>	99	1/158	1/15800	1/998560000	1/63200
Glycogen storage disease IV	AR	<i>GBE1</i>	92	1/387	1/4838	1/93605625	1/19350
N-acetylglutamate synthase deficiency	AR	<i>NAGS</i>	99	<1/500	1/50000	1/10000000000	1/200000
Peroxisome biogenesis disorder 6A (Zellweger)	AR	<i>PEX10</i>	86	1/500	1/3571	1/51020408	1/14286
Peroxisome biogenesis disorder 3A (Zellweger)	AR	<i>PEX12</i>	99	1/373	1/37300	1/5565160000	1/149200
Peroxisome biogenesis disorder 5A (Zellweger)	AR	<i>PEX2</i>	99	1/500	1/50000	1/10000000000	1/200000
Combined oxidative phosphorylation deficiency 3	AR	<i>TSFM</i>	99	<1/500	1/50000	1/10000000000	1/200000
Adrenal hypoplasia, congenital	XL	<i>NR0B1</i>	99	1/12500	1/1250000	1/6250000000000	1/5000000
Emphysema due to AAT deficiency	AR	<i>SERPINA1</i>	99	1/33	1/3300	1/43560000	1/13200
Mannosidosis, alpha-, types I and II	AR	<i>MAN2B1</i>	99	1/354	1/35400	1/5012640000	1/141600
Alpha-methylacetoacetic aciduria	AR	<i>ACAT1</i>	89	<1/500	1/4545	1/82644628	1/18182
Carbamoylphosphate synthetase I deficiency	AR	<i>CPS1</i>	99	1/570	1/57000	1/12996000000	1/228000
Galactosemia	AR	<i>GALT</i>	99	1/110	1/11000	1/484000000	1/44000
Galactose epimerase deficiency	AR	<i>GALE</i>	99	1/100	1/10000	1/400000000	1/40000
Galactokinase deficiency with cataracts	AR	<i>GALK1</i>	99	1/110	1/11000	1/484000000	1/44000
Galactosemia IV	AR	<i>GALM</i>	99	1/250	1/25000	1/2500000000	1/100000
Pyruvate carboxylase deficiency	AR	<i>PC</i>	99	1/250	1/25000	1/2500000000	1/100000
Phenylketonuria	AR	<i>PAH</i>	99	1/99	1/9900	1/392040000	1/39600
Propionicacidemia	AR	<i>PCCB</i>	96	1/224	1/5600	1/125440000	1/22400

Propionic acidemia	AR	PCCA	82	1/224	1/1244	1/6194568	1/4978
Pyruvate dehydrogenase E1-beta deficiency	AR	PDHB	96	<1/500	1/12500	1/625000000	1/50000
Cerebral creatine deficiency syndrome 1	XL	SLC6A8	94	<1/500	1/8333	1/277777778	1/33333
Cholesteryl ester storage disease	AR	LIPA	99	<1/500	1/50000	1/10000000000	1/200000
Hypophosphatasia, infantile	AR	ALPL	99	1/158	1/15800	1/998560000	1/63200
Butyrylcholinesterase deficiency	AR	BCHE	99	1/28	1/2800	1/31360000	1/11200
Multiple sulfatase deficiency	AR	SUMF1	99	1/500	1/50000	1/10000000000	1/200000
Dihydropyrimidine dehydrogenase deficiency	AR	DPYD	99	<1/500	1/50000	1/10000000000	1/200000
Fabry disease	XL	GLA	93	<1/500	1/7143	1/204081633	1/28571
Aromatase deficiency	AR	CYP19A1	99	<1/500	1/50000	1/10000000000	1/200000
Hyperphenylalaninemia, mild, non-BH4-deficient	AR	DNAJC12	94	<1/200	1/3333	1/444444444	1/13333
Glycine encephalopathy	AR	GLDC	99	1/193	1/19300	1/1489960000	1/77200
Maple syrup urine disease, type Ia	AR	BCKDHA	99	1/321	1/32100	1/4121640000	1/128400
Maple syrup urine disease, type Ib	AR	BCKDHB	90	1/364	1/3640	1/52998400	1/14560
Dihydroipoamide dehydrogenase deficiency	AR	DLD	99	1/500	1/50000	1/10000000000	1/200000
Maple syrup urine disease, type II	AR	DBT	69	1/481	1/1552	1/9630010	1/6206
Wilson disease	AR	ATP7B	99	1/90	1/9000	1/324000000	1/36000
Mitochondrial DNA depletion syndrome 6 (hepatocerebral type)	AR	MPV17	99	<1/500	1/50000	1/10000000000	1/200000
Methionine adenosyltransferase deficiency, autosomal recessive	AR/AD	MAT1A	99	3/500	1/16667	1/1111111111	1/66667
Homocystinuria-megaloblastic anemia, cbl E type	AR	MTRR	93	<1/500	1/7143	1/204081633	1/28571
Homocystinuria-megaloblastic anemia, cblG complementation type	AR	MTR	94	<1/2000	1/33333	1/4444444444	1/133333
Homocystinuria due to MTHFR deficiency	AR	MTHFR	99	1/2	1/200	1/160000	1/800
Hyperornithinemia-hyperammonemia-homocitrullinemia syndrome	AR	SLC25A15	92	<1/500	1/6250	1/156250000	1/25000
Thrombosis, hyperhomocysteinemic	AR	CBS	10	1/224	1/249	1/247783	1/996
Gaucher disease	AR	GBA1	64	3/500	1/463	1/857339	1/1852
Sitosterolemia 1	AR	ABCG8	99	9/1000	1/11111	1/493827160	1/44444
Sitosterolemia 2	AR	ABCG5	93	9/1000	1/1587	1/10078105	1/6349
Citrullinemia, type II, neonatal-onset	AR	SLC25A13	99	1/65	1/6500	1/169000000	1/26000
Citrullinemia	AR	ASS1	41	1/119	1/202	1/162723	1/807
Cerebral creatine deficiency syndrome 2	AR	GAMT	72	1/371	1/1325	1/7022500	1/5300
Cystinosis, ocular nonnephropathic	AR	CTNS	93	1/158	1/2257	1/20378776	1/9029
Peroxisomal acyl-CoA oxidase deficiency	AR	ACOX1	99	<1/500	1/50000	1/10000000000	1/200000
VLCAD deficiency	AR	ACADVL	99	1/200	1/20000	1/1600000000	1/80000
Lipoprotein lipase deficiency	AR	LPL	99	1/500	1/50000	1/10000000000	1/200000
Methylmalonyl-CoA epimerase deficiency	AR	MCEE	99	3/500	1/16667	1/1111111111	1/66667
Methylmalonic aciduria, vitamin B12-responsive, cblA type	AR	MMAA	99	1/301	1/30100	1/3624040000	1/120400
Methylmalonic aciduria, vitamin B12-responsive, cblB type	AR	MMAB	99	1/435	1/43500	1/7569000000	1/174000
Methylmalonic aciduria and homocystinuria, cblF type	AR	LMBRD1	90	1/25	1/250	1/250000	1/1000
Mental retardation, X-linked 3 (methylmalonic acidemia and homocystinemia, cblX type)	XL	HCFC1	86	1/25	1/179	1/127551	1/714
Methylmalonic aciduria, mut(0) type	AR	MMUT	96	<9/1000	1/2778	1/30864198	1/11111
Methylmalonic aciduria and homocystinuria, cblC type	AR	MMACHC	99	1/134	1/13400	1/718240000	1/53600
Methylmalonic aciduria and homocystinuria, cblC type, digenic	AR	PRDX1	99	1/25	1/2500	1/25000000	1/10000
Methylmalonic aciduria and homocystinuria, cblD type	AR	MMADHC	83	<1/500	1/2941	1/34602076	1/11765
Argininosuccinic aciduria	AR	ASL	97	1/200	1/6667	1/177777778	1/26667
Argininemia	AR	ARG1	98	<4/1000	1/12500	1/625000000	1/50000
Lysinuric protein intolerance	AR	SLC7A7	99	<1/125	1/12500	1/625000000	1/50000
Tyrosinemia, type III	AR	HPD	99	<1/500	1/50000	1/10000000000	1/200000
Tyrosinemia, type II	AR	TAT	99	1/200	1/20000	1/1600000000	1/80000
Tyrosinemia, type I	AR	FAH	99	1/99	1/9900	1/392040000	1/39600

Lipoid adrenal hyperplasia	AR	<i>STAR</i>	99	<1/500	1/50000	1/10000000000	1/200000
Combined malonic and methylmalonic aciduria	AR	<i>ACSF3</i>	99	<1/500	1/50000	1/10000000000	1/200000
Combined oxidative phosphorylation deficiency 1	AR	<i>GFM1</i>	96	<1/500	1/12500	1/625000000	1/50000
Phosphoglycerate dehydrogenase deficiency	AR	<i>PHGDH</i>	99	<1/500	1/50000	1/10000000000	1/200000
Menkes disease	XL	<i>ATP7A</i>	93	<1/500	1/7143	1/204081633	1/28571
Molybdenum cofactor deficiency A	AR	<i>MOCS1</i>	88	<3/500	1/1389	1/7716049	1/5556
Cerebrotendinous xanthomatosis	AR	<i>CYP27A1</i>	99	1/500	1/50000	1/10000000000	1/200000
Niemann-Pick disease, type B	AR	<i>SMPD1</i>	98	<9/1000	1/5556	1/123456790	1/22222
Niemann-Pick disease, type C1	AR	<i>NPC1</i>	99	<3/500	1/16667	1/1111111111	1/66667
Niemann-pick disease, type C2	AR	<i>NPC2</i>	99	<3/500	1/16667	1/1111111111	1/66667
Mucopolysaccharidosis type IIIC (Sanfilippo C)	AR	<i>HGSNAT</i>	96	1/434	1/10850	1/470890000	1/43400
Mucopolysaccharidosis Ih	AR	<i>IDUA</i>	67	<1/500	1/1515	1/9182736	1/6061
Mucopolysaccharidosis type IIIA (Sanfilippo A)	AR	<i>SGSH</i>	98	1/454	1/22700	1/2061160000	1/90800
Mucopolysaccharidosis type IIIB (Sanfilippo B)	AR	<i>NAGLU</i>	89	<1/500	1/4545	1/82644628	1/18182
Mucopolysaccharidosis II	XL	<i>IDS</i>	87	<1/500	1/3846	1/59171598	1/15385
Mucopolysaccharidosis IVA	AR	<i>GALNS</i>	96	1/224	1/5600	1/125440000	1/22400
Mucopolysaccharidosis type IVB (Morquio)	AR	<i>GLB1</i>	99	1/134	1/13400	1/718240000	1/53600
Mucopolysaccharidosis type IX	AR	<i>HYAL1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Mucopolysaccharidosis VII	AR	<i>GUSB</i>	95	1/250	1/5000	1/100000000	1/20000
Mucopolysaccharidosis type VI (Maroteaux-Lamy)	AR	<i>ARSB</i>	92	1/250	1/3125	1/39062500	1/12500
Mucopolipidosis II alpha;beta	AR	<i>GNPTAB</i>	98	<1/500	1/25000	1/2500000000	1/100000
Mucopolipidosis IV	AR	<i>MCOLN1</i>	94	1/300	1/5000	1/100000000	1/20000
Ornithine transcarbamylase deficiency	XL	<i>OTC</i>	99	<1/500	1/50000	1/10000000000	1/200000
Hypoaldosteronism, congenital, due to CMO I deficiency	AR	<i>CYP11B2</i>	99	<1/500	1/50000	1/10000000000	1/200000
Holocarboxylase synthetase deficiency	AR	<i>HLCS</i>	99	1/500	1/50000	1/10000000000	1/200000
Carnitine-acylcarnitine translocase deficiency	AR	<i>SLC25A20</i>	98	<1/500	1/25000	1/2500000000	1/100000
CPT II deficiency, myopathic, stress-induced	AR/AD	<i>CPT2</i>	99	<1/500	1/50000	1/10000000000	1/200000
CPT deficiency, hepatic, type IA	AR	<i>CPT1A</i>	98	1/354	1/17700	1/1253160000	1/70800
Trifunctional protein deficiency	AR	<i>HADHB</i>	74	1/500	1/1923	1/14792899	1/7692
Biotinidase deficiency	AR	<i>BTBD</i>	99	1/125	1/12500	1/625000000	1/50000
Glycogen storage disease II	AR	<i>GAA</i>	37	1/100	1/159	1/100781	1/635
Glycogen storage disease Ia	AR	<i>G6PC1</i>	99	1/177	1/17700	1/1253160000	1/70800
Glycogen storage disease Ib	AR	<i>SLC37A4</i>	97	1/158	1/5267	1/110951111	1/21067
Glycogen storage disease VII	AR	<i>PFKM</i>	99	<1/500	1/50000	1/10000000000	1/200000
McArdle disease	AR	<i>PYGM</i>	99	<1/500	1/50000	1/10000000000	1/200000
Aspartylglucosaminuria	AR	<i>AGA</i>	99	<1/500	1/50000	1/10000000000	1/200000
Asparagine synthetase deficiency	AR	<i>ASNS</i>	53	<1/500	1/1064	1/4526935	1/4255
Glutaricaciduria, type I	AR	<i>GCDH</i>	99	1/87	1/8700	1/302760000	1/34800
Glutaric acidemia IIA	AR	<i>ETFA</i>	99	1/500	1/50000	1/10000000000	1/200000
Glutaric acidemia IIB	AR	<i>ETFB</i>	99	1/500	1/50000	1/10000000000	1/200000
Glutaric acidemia IIC	AR	<i>ETFDH</i>	99	1/500	1/50000	1/10000000000	1/200000
Bile acid synthesis defect, congenital, 1	AR	<i>HSD3B7</i>	61	1/500	1/1282	1/6574622	1/5128
Bile acid synthesis defect, congenital, 2	AR	<i>AKR1D1</i>	99	<3/500	1/16667	1/1111111111	1/66667
Bile acid synthesis defect, congenital, 3	AR	<i>CYP7B1</i>	94	<1/200	1/3333	1/44444444	1/13333
Bile acid synthesis defect, congenital, 4	AR	<i>AMACR</i>	99	<1/200	1/20000	1/1600000000	1/80000
Bile acid synthesis defect, congenital, 5	AR	<i>ABCD3</i>	99	<1/200	1/20000	1/1600000000	1/80000
Bile acid synthesis defect, congenital, 6	AR	<i>ACOX2</i>	99	<1/200	1/20000	1/1600000000	1/80000
Hyperbilirubinemia, familial transient neonatal	AR	<i>UGT1A1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type IIb	AR	<i>MOGS</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type IIc	AR	<i>SLC35C1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type IId	AR	<i>B4GALT1</i>	99	<1/500	1/50000	1/10000000000	1/200000

Congenital disorder of glycosylation, type If	AR	<i>SLC35A1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type Im	AR	<i>DOLK</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type Ia	AR	<i>PMM2</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type Ib	AR	<i>MPI</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type Ic	AR	<i>ALG6</i>	63	<1/500	1/1351	1/7304602	1/5405
Congenital disorder of glycosylation, type Id	AR	<i>ALG3</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type Ie	AR	<i>DPM1</i>	92	<1/500	1/6250	1/156250000	1/25000
Congenital disorder of glycosylation, type If	AR	<i>MPDU1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type Ig	AR	<i>ALG12</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type Ih	AR	<i>ALG8</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type Ij	AR	<i>DPAGT1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type Ik	AR	<i>ALG1</i>	25	<1/500	1/667	1/1777778	1/2667
Congenital disorder of glycosylation, type In	AR	<i>RFT1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Congenital disorder of glycosylation, type Iq	AR	<i>SRD5A3</i>	99	<1/500	1/50000	1/10000000000	1/200000
Mitochondrial complex I deficiency, nuclear type 20	AR	<i>ACAD9</i>	99	<1/500	1/50000	1/10000000000	1/200000
Mitochondrial complex I deficiency, nuclear type 16	AR	<i>NDUFAF5</i>	67	1/447	1/1355	1/7339174	1/5418
Mitochondrial complex I deficiency, nuclear type 9	AR	<i>NDUFS6</i>	99	<1/500	1/50000	1/10000000000	1/200000
Mitochondrial trifunctional protein deficiency	AR	<i>HADHA</i>	93	3/200	1/952	1/3628118	1/3810
Adenosine deaminase deficiency, partial	AR	<i>ADA</i>	98	1/224	1/11200	1/501760000	1/44800
Fumarase deficiency	AR	<i>FH</i>	88	<1/500	1/4167	1/69444444	1/16667
Fructose intolerance, hereditary	AR	<i>ALDOB</i>	99	1/122	1/12200	1/595360000	1/48800
Ethylmalonic encephalopathy	AR	<i>ETHE1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Isobutyryl-CoA dehydrogenase deficiency	AR	<i>ACAD8</i>	99	1/125	1/12500	1/625000000	1/50000
Isovaleric acidemia	AR	<i>IVD</i>	99	1/167	1/16700	1/1115560000	1/66800
Hyperoxaluria, primary, type II	AR	<i>GRHPR</i>	95	<1/500	1/10000	1/400000000	1/40000
Hyperoxaluria, primary, type III	AR	<i>HOGA1</i>	99	1/184	1/18400	1/1354240000	1/73600
Hyperoxaluria, primary, type 1	AR	<i>AGXT</i>	99	1/120	1/12000	1/576000000	1/48000
Carnitine deficiency, systemic primary	AR	<i>SLC22A5</i>	99	1/129	1/12900	1/665640000	1/51600
Mucopolidosis III gamma	AR	<i>GNPTG</i>	94	<1/500	1/8333	1/277777778	1/33333
Mucopolysaccharidosis type IIID	AR	<i>GNS</i>	93	1/500	1/7143	1/204081633	1/28571
Acyl-CoA dehydrogenase, medium chain, deficiency of	AR	<i>ACADM</i>	99	<3/500	1/16667	1/1111111111	1/66667
Acyl-CoA dehydrogenase, short-chain, deficiency of	AR	<i>ACADS</i>	99	1/85	1/8500	1/289000000	1/34000

Musculoskeletal System

Achondrogenesis Ib	AR	<i>SLC26A2</i>	99	1/158	1/15800	1/998560000	1/63200
Myasthenic syndrome, congenital	AR	<i>CHRNE</i>	99	1/408	1/40800	1/6658560000	1/163200
Emery-Dreifuss muscular dystrophy 1, X-linked	XL	<i>EMD</i>	87	<1/500	1/3846	1/59171598	1/15385
Emery-Dreifuss muscular dystrophy 3, autosomal recessive	AR	<i>LMNA</i>	93	<1/500	1/7143	1/204081633	1/28571
Emery-Dreifuss muscular dystrophy 6, X-linked	XL	<i>FHL1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Muscular dystrophy-dystroglycanopathy	AR	<i>FKTN</i>	99	<1/500	1/50000	1/10000000000	1/200000
Muscular dystrophy, limb-girdle, autosomal recessive 23	AR	<i>LAMA2</i>	99	<1/500	1/50000	1/10000000000	1/200000
Schimke immunoskeletal dysplasia	AR	<i>SMARCA1</i>	99	1/500	1/50000	1/10000000000	1/200000
Nonaka myopathy	AR	<i>GNE</i>	99	<1/500	1/50000	1/10000000000	1/200000
Osteopetrosis, autosomal recessive 1	AR	<i>TCIRG1</i>	80	1/250	1/1250	1/6250000	1/5000
Osteopetrosis, autosomal recessive 4	AR	<i>CLCN7</i>	93	1/250	1/3571	1/51020408	1/14286
Duchenne muscular dystrophy	XL	<i>DMD</i>	80	<1/500	1/2500	1/25000000	1/10000
Nemaline myopathy 2, autosomal recessive	AR	<i>NEB</i>	94	1/112	1/1867	1/13937778	1/7467
Muscular dystrophy-dystroglycanopathy (congenital with brain and eye anomalies), type A, 6	AR	<i>LARGE1</i>	99	1/125	1/12500	1/625000000	1/50000
Spinal muscular atrophy-2	AR	<i>SMN1</i>	10	1/50	1/56	1/12346	1/222
Spinal and bulbar muscular atrophy of Kennedy	XL	<i>AR</i>	54	<1/500	1/1087	1/4725898	1/4348
Spondylocostal dysostosis 2, autosomal recessive	AR	<i>MESP2</i>	99	<1/500	1/50000	1/10000000000	1/200000
Carpenter syndrome	AR	<i>RAB23</i>	95	<1/500	1/10000	1/400000000	1/40000

Congenital arthrogryposis with anterior horn cell disease	AR	<i>GLE1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Myasthenic syndrome, congenital, 11, associated with acetylcholine receptor deficiency	AR	<i>RAPSN</i>	99	<1/500	1/50000	1/10000000000	1/200000
Myasthenic syndrome, congenital, 10	AR	<i>DOK7</i>	63	1/163	1/441	1/776304	1/1762
Myasthenic syndrome, congenital, 1B, fast-channel	AR/AD	<i>CHRNA1</i>	99	1/577	1/57700	1/13317160000	1/230800
Myasthenic syndrome, congenital, 5	AR	<i>COLQ</i>	99	1/163	1/16300	1/1062760000	1/65200
Myasthenic syndrome, congenital, 6, presynaptic	AR	<i>CHAT</i>	99	1/272	1/27200	1/2959360000	1/108800
Charcot-Marie-Tooth disease, type 4D	AR	<i>NDRG1</i>	99	1/22	1/2200	1/19360000	1/8800
Muscular dystrophy, limb-girdle, autosomal recessive 8	AR	<i>TRIM32</i>	99	<1/500	1/50000	1/10000000000	1/200000
Muscular dystrophy-dystroglycanopathy (limb-girdle), type C, 1	AR	<i>POMT1</i>	99	1/843	1/84300	1/28425960000	1/337200
Muscular dystrophy, limb-girdle, autosomal recessive 12	AR	<i>ANO5</i>	97	<1/500	1/16667	1/1111111111	1/66667
Muscular dystrophy-dystroglycanopathy (limb-girdle), type C, 2	AR	<i>POMT2</i>	99	<1/500	1/50000	1/10000000000	1/200000
Muscular dystrophy-dystroglycanopathy (limb-girdle), type C, 9	AR	<i>DAG1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Muscular dystrophy-dystroglycanopathy (limb-girdle), type C, 14	AR	<i>GMPPB</i>	99	<1/500	1/50000	1/10000000000	1/200000
Muscular dystrophy, limb-girdle, autosomal recessive 1	AR	<i>CAPN3</i>	99	<1/500	1/50000	1/10000000000	1/200000
Muscular dystrophy-dystroglycanopathy (congenital with brain and eye anomalies), type A, 3	AR	<i>POMGNT1</i>	99	1/462	1/46200	1/8537760000	1/184800
Muscular dystrophy, limb-girdle, autosomal recessive 2	AR	<i>DYSF</i>	99	<1/500	1/50000	1/10000000000	1/200000
Muscular dystrophy, limb-girdle, autosomal recessive 3	AR	<i>SGCA</i>	98	<1/500	1/25000	1/2500000000	1/100000
Muscular dystrophy, limb-girdle, autosomal recessive 4	AR	<i>SGCB</i>	78	1/500	1/2273	1/20661157	1/9091
Muscular dystrophy, limb-girdle, autosomal recessive 5	AR	<i>SGCG</i>	99	1/381	1/38100	1/5806440000	1/152400
Muscular dystrophy, limb-girdle, autosomal recessive 6	AR	<i>SGCD</i>	74	<1/500	1/1923	1/14792899	1/7692
Muscular dystrophy-dystroglycanopathy (limb-girdle), type C, 5	AR	<i>FKRP</i>	95	<1/500	1/10000	1/400000000	1/40000
Rhizomelic chondrodysplasia punctata, type 1	AR	<i>PEX7</i>	66	1/158	1/465	1/863806	1/1859
Rhizomelic chondrodysplasia punctata, type 3	AR	<i>AGPS</i>	99	<1/500	1/50000	1/10000000000	1/200000
Chondrodysplasia punctata, X-linked recessive	XL	<i>ARSL</i>	99	<1/500	1/50000	1/10000000000	1/200000
Myotubular myopathy, X-linked	XL	<i>MTM1</i>	83	<1/500	1/2941	1/34602076	1/11765
Pycnodysostosis	AR	<i>CTSK</i>	99	<1/500	1/50000	1/10000000000	1/200000

Respiratory System

Ciliary dyskinesia, primary, 1, with or without situs inversus	AR	<i>DNAI1</i>	99	1/230	1/23000	1/2116000000	1/92000
Ciliary dyskinesia, primary, 9, with or without situs inversus	AR	<i>DNAI2</i>	99	1/447	1/44700	1/7992360000	1/178800
Ciliary dyskinesia, primary, 16	AR	<i>DNAL1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Ciliary dyskinesia, primary, 3, with or without situs inversus	AR	<i>DNAH5</i>	99	1/142	1/14200	1/806560000	1/56800
Ciliary dyskinesia, primary, 17	AR	<i>DNAAF19</i>	99	1/316	1/31600	1/3994240000	1/126400
Ciliary dyskinesia, primary, 30	AR	<i>ODAD3</i>	99	1/365	1/36500	1/5329000000	1/146000
Ciliary dyskinesia, primary, 14	AR	<i>CCDC39</i>	62	1/211	1/555	1/1233269	1/2221

Urinary System

Alkaptonuria	AR	<i>HGD</i>	99	<1/500	1/50000	1/10000000000	1/200000
Polycystic kidney disease 5	AR	<i>DZIP1L</i>	99	<1/100	1/10000	1/400000000	1/40000
Polycystic kidney disease 4, with or without hepatic disease	AR	<i>PKHD1</i>	99	<1/100	1/10000	1/400000000	1/40000
Nephrotic syndrome, type 1	AR	<i>NPHS1</i>	98	1/289	1/14450	1/835210000	1/57800
Nephrotic syndrome, type 2	AR	<i>NPHS2</i>	99	3/500	1/16667	1/1111111111	1/66667
Nephrotic syndrome, type 3	AR	<i>PLCE1</i>	99	3/500	1/16667	1/1111111111	1/66667
Nephrotic syndrome, type 6	AR	<i>PTPRO</i>	99	3/500	1/16667	1/1111111111	1/66667
Alport syndrome 2, autosomal recessive	AR	<i>COL4A3</i>	99	1/267	1/26700	1/2851560000	1/106800
Alport syndrome 2, autosomal recessive	AR	<i>COL4A4</i>	64	1/267	1/742	1/2200278	1/2967
Dent disease 2	XL	<i>OCRL</i>	95	<1/500	1/10000	1/400000000	1/40000
Diabetes insipidus, nephrogenic	AR/AD	<i>AQP2</i>	89	<1/500	1/4545	1/82644628	1/18182
Nephronophthisis 3	AR	<i>NPHP3</i>	71	<3/1000	1/1149	1/5284714	1/4598

Immune System

Chronic granulomatous disease 2, autosomal recessive	AR	<i>NCF2</i>	99	1/224	1/22400	1/2007040000	1/89600
Chronic granulomatous disease 4, autosomal recessive	AR	<i>CYBA</i>	89	1/224	1/2036	1/16587107	1/8145
Chronic granulomatous disease 5, autosomal recessive	AR	<i>CYBC1</i>	99	1/224	1/22400	1/2007040000	1/89600
Chronic granulomatous disease, X-linked	XL	<i>CYBB</i>	84	<1/500	1/3125	1/39062500	1/12500
Familial Mediterranean fever, AR	AR	<i>MEFV</i>	99	1/20	1/2000	1/16000000	1/8000
Wiskott-Aldrich syndrome	XL	<i>WAS</i>	27	<1/500	1/685	1/1876525	1/2740
Agammaglobulinemia, X-linked 1	XL	<i>BTK</i>	99	<1/500	1/50000	1/10000000000	1/200000
Lymphoproliferative syndrome, X-linked, 2	XL	<i>XIAP</i>	80	<1/500	1/2500	1/25000000	1/10000
Lymphoproliferative syndrome, X-linked, 1	XL	<i>SH2D1A</i>	93	<1/500	1/7143	1/204081633	1/28571
Combined cellular and humoral immune defects with granulomas	AR	<i>RAG1</i>	99	1/137	1/13700	1/750760000	1/54800
Combined cellular and humoral immune defects with granulomas	AR	<i>RAG2</i>	99	1/137	1/13700	1/750760000	1/54800
Severe combined immunodeficiency, X-linked	XL	<i>IL2RG</i>	99	<1/500	1/50000	1/10000000000	1/200000
Severe combined immunodeficiency, Athabaskan type	AR	<i>DCLRE1C</i>	99	<1/500	1/50000	1/10000000000	1/200000
Immunodeficiency, X-linked, with hyper-IgM	XL	<i>CD40LG</i>	97	<1/500	1/16667	1/1111111111	1/66667
Bare lymphocyte syndrome, type II, complementation group A	AR	<i>CIITA</i>	99	<1/500	1/50000	1/10000000000	1/200000
Hemophagocytic lymphohistiocytosis, familial, 2	AR	<i>PRF1</i>	99	3/500	1/16667	1/1111111111	1/66667
Hemophagocytic lymphohistiocytosis, familial, 3	AR	<i>UNC13D</i>	99	3/500	1/16667	1/1111111111	1/66667
Hemophagocytic lymphohistiocytosis, familial, 4	AR	<i>STX11</i>	99	3/500	1/16667	1/1111111111	1/66667
Hemophagocytic lymphohistiocytosis, familial, 5	AR	<i>STXBP2</i>	99	3/500	1/16667	1/1111111111	1/66667
Endocrine System							
Hyperinsulinemic hypoglycemia, familial, 1	AR/AD	<i>ABCC8</i>	99	1/112	1/11200	1/501760000	1/44800
Hyperinsulinemic hypoglycemia, familial, 2	AR/AD	<i>KCNJ11</i>	99	1/423	1/42300	1/7157160000	1/169200
Hyperinsulinemic hypoglycemia, familial, 4	AR	<i>HADH</i>	92	<1/500	1/6250	1/156250000	1/25000
Autoimmune polyendocrinopathy syndrome , type I, with or without reversible metaphyseal dysplasia	AR/AD	<i>AIRE</i>	99	1/150	1/15000	1/900000000	1/60000
Pituitary hormone deficiency, combined, 2	AR	<i>PROP1</i>	99	1/45	1/4500	1/81000000	1/18000
Hypothyroidism, congenital, nongoitrous, 1	AR	<i>TSHR</i>	99	1/50	1/5000	1/100000000	1/20000
Hypothyroidism, congenital, nongoitrous 4	AR	<i>TSHB</i>	99	1/50	1/5000	1/100000000	1/20000
Pituitary hormone deficiency, combined, 3	AR	<i>LHX3</i>	99	1/45	1/4500	1/81000000	1/18000
Pituitary hormone deficiency, combined, 1	AR/AD	<i>POU1F1</i>	94	1/267	1/4450	1/79210000	1/17800
Hypercholesterolemia, familial, 4	AR	<i>LDLRAP1</i>	99	1/8	1/800	1/2560000	1/3200
Integumentary System							
Ehlers-Danlos syndrome, dermatosparaxis type	AR	<i>ADAMTS2</i>	90	<1/500	1/5000	1/100000000	1/20000
Ocular albinism, type I, NettleSHIP-Falls type	XL	<i>GPR143</i>	83	1/25000	1/147059	1/86505190311	1/588235
Acrodermatitis enteropathica	AR	<i>SLC39A4</i>	99	<1/500	1/50000	1/10000000000	1/200000
Ichthyosis, congenital, autosomal recessive 4A	AR	<i>ABCA12</i>	99	1/284	1/28400	1/3226240000	1/113600
Epidermolysis bullosa, junctional, localisata variant	AR	<i>COL17A1</i>	96	<1/2000	1/50000	1/10000000000	1/200000
Ataxia-telangiectasia	AR	<i>ATM</i>	89	1/100	1/909	1/3305785	1/3636
Epidermolysis bullosa	AR	<i>LAMA3</i>	99	1/781	1/78100	1/24398440000	1/312400
Epidermolysis bullosa, junctional, Herlitz type	AR	<i>LAMB3</i>	99	1/781	1/78100	1/24398440000	1/312400
Epidermolysis bullosa, junctional, Herlitz type	AR	<i>LAMC2</i>	99	1/781	1/78100	1/24398440000	1/312400
Epidermolysis bullosa dystrophica	AR	<i>COL7A1</i>	99	1/196	1/19600	1/1536640000	1/78400
Dyskeratosis congenita, autosomal recessive 5	AR/AD	<i>RTEL1</i>	98	1/500	1/25000	1/2500000000	1/100000
Ichthyosis, congenital, autosomal recessive 1	AR	<i>TGM1</i>	99	1/224	1/22400	1/2007040000	1/89600
Albinism, oculocutaneous, type IB	AR	<i>TYR</i>	99	3/200	1/6667	1/177777778	1/26667
Albinism, oculocutaneous, type II, modifier of	AR	<i>MC1R</i>	99	3/200	1/6667	1/177777778	1/26667
Albinism, oculocutaneous, type III	AR	<i>TYRP1</i>	99	3/200	1/6667	1/177777778	1/26667
Albinism, oculocutaneous, type IV	AR	<i>SLC45A2</i>	99	3/200	1/6667	1/177777778	1/26667
Skin;;hair;;eye pigmentation 4, fair	AR	<i>SLC24A5</i>	38	3/200	1/108	1/46248	1/430
Albinism, oculocutaneous, type VII	AR	<i>LRMDA</i>	99	3/200	1/6667	1/177777778	1/26667

Oculocutaneous albinism, type VIII	AR	<i>DCT</i>	99	3/200	1/6667	1/177777778	1/26667
Xeroderma pigmentosum, group A	AR	<i>XPA</i>	69	1/500	1/1613	1/10405827	1/6452
Xeroderma pigmentosum, group C	AR	<i>XPC</i>	99	1/500	1/50000	1/10000000000	1/200000
Albinism, brown oculocutaneous	AR	<i>OCA2</i>	99	3/200	1/6667	1/177777778	1/26667
Nervous System							
Salt and pepper developmental regression syndrome	AR	<i>ST3GAL5</i>	92	9/50	1/69	1/19290	1/278
Ceroid lipofuscinosis, neuronal, 3	AR	<i>CLN3</i>	93	1/230	1/3286	1/43183673	1/13143
CRASH syndrome Corpus callosum, partial agenesis of;;CRASH syndrome;;Hydrocephalus due to aqueductal stenosis;;Hydrocephalus with congenital idiopathic intestinal pseudoobstruction;;Hydrocephalus with Hirschsprung disease;;MASA syndrome	XL	<i>L1CAM</i>	99	<1/500	1/50000	1/10000000000	1/200000
Segawa syndrome, recessive	AR	<i>TH</i>	97	1/224	1/7467	1/223004444	1/29867
Tay-Sachs disease	AR	<i>HEXA</i>	99	1/300	1/30000	1/3600000000	1/120000
Lissencephaly, X-linked	XL	<i>DCX</i>	99	<1/500	1/50000	1/10000000000	1/200000
Leukoencephalopathy with vanishing white matter	AR	<i>EIF2B5</i>	99	<1/500	1/50000	1/10000000000	1/200000
Spastic ataxia, Charlevoix-Saguenay type	AR	<i>SACS</i>	99	<1/500	1/50000	1/10000000000	1/200000
Ataxia with isolated vitamin E deficiency	AR	<i>TTPA</i>	83	<1/500	1/2941	1/34602076	1/11765
Arthrogryposis, mental retardation, and seizures	AR	<i>SLC35A3</i>	41	<1/500	1/847	1/2872738	1/3390
Dysautonomia, familial	AR	<i>ELP1</i>	99	1/300	1/30000	1/3600000000	1/120000
Microcephaly, postnatal progressive, with seizures and brain atrophy	AR	<i>MED17</i>	99	<1/500	1/50000	1/10000000000	1/200000
Spastic paraplegia 15, autosomal recessive	AR	<i>ZFYVE26</i>	99	<1/500	1/50000	1/10000000000	1/200000
Spastic paraplegia 49, autosomal recessive	AR	<i>TECPR2</i>	99	<1/500	1/50000	1/10000000000	1/200000
Megalencephalic leukoencephalopathy with subcortical cysts	AR	<i>MLC1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Canavan disease	AR	<i>ASPA</i>	99	1/57	1/5700	1/129960000	1/22800
Hydrolethalus syndrome	AR	<i>HYLS1</i>	99	7/500	1/7143	1/204081633	1/28571
Pontocerebellar hypoplasia type 1A	AR	<i>VRK1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Pontocerebellar hypoplasia, type 1B	AR	<i>EXOSC3</i>	77	<1/500	1/2174	1/18903592	1/8696
Pontocerebellar hypoplasia type 2D	AR	<i>SEPSECS</i>	89	<1/500	1/4545	1/82644628	1/18182
Pontocerebellar hypoplasia, type 6	AR	<i>RARS2</i>	97	<1/500	1/16667	1/1111111111	1/66667
Agenesis of the corpus callosum with peripheral neuropathy	AR	<i>SLC12A6</i>	99	<1/500	1/50000	1/10000000000	1/200000
Sandhoff disease, infantile, juvenile, and adult forms	AR	<i>HEXB</i>	97	1/600	1/20000	1/1600000000	1/80000
Ceroid lipofuscinosis, neuronal, 10	AR	<i>CTSD</i>	99	1/1761	1/176100	1/12404484000	1/704400
Ceroid lipofuscinosis, neuronal, 1	AR	<i>PPT1</i>	99	1/368	1/36800	1/5416960000	1/147200
Ceroid lipofuscinosis, neuronal, 2	AR	<i>TPP1</i>	99	1/252	1/25200	1/2540160000	1/100800
Ceroid lipofuscinosis, neuronal, 5	AR	<i>CLN5</i>	66	<1/500	1/1471	1/8650519	1/5882
Ceroid lipofuscinosis, neuronal, 6	AR	<i>CLN6</i>	86	<1/500	1/3571	1/51020408	1/14286
Ceroid lipofuscinosis, neuronal, 7	AR	<i>MFSD8</i>	95	<1/500	1/10000	1/400000000	1/40000
Ceroid lipofuscinosis, neuronal, 8, Northern epilepsy variant	AR	<i>CLN8</i>	99	<1/500	1/50000	1/10000000000	1/200000
Adrenoleukodystrophy	XL	<i>ABCD1</i>	86	1/100	1/714	1/2040816	1/2857
Polymicrogyria, bilateral frontoparietal	AR	<i>ADGRG1</i>	98	<1/500	1/25000	1/2500000000	1/100000
Choreoacanthocytosis	AR	<i>VPS13A</i>	80	<1/500	1/2500	1/25000000	1/10000
Mitochondrial complex IV deficiency, nuclear type 5, (French-Canadian)	AR	<i>LRPPRC</i>	91	1/447	1/4967	1/98671111	1/19867
Metachromatic leukodystrophy	AR	<i>ARSA</i>	99	1/100	1/10000	1/400000000	1/40000
Infantile neuroaxonal dystrophy 1	AR	<i>PLA2G6</i>	97	1/500	1/16667	1/1111111111	1/66667
Krabbe disease	AR	<i>GALC</i>	97	1/158	1/5267	1/110951111	1/21067
Metachromatic leukodystrophy due to SAP-b deficiency	AR	<i>PSAP</i>	99	<1/500	1/50000	1/10000000000	1/200000
Visual System							
Achromatopsia 3	AR	<i>CNGB3</i>	99	1/87	1/8700	1/302760000	1/34800
Progressive external ophthalmoplegia, autosomal recessive 1;;Mitochondrial DNA depletion syndrome 4	AR	<i>POLG</i>	99	1/113	1/11300	1/510760000	1/45200
Glaucoma 3A, primary open angle, congenital, juvenile, or adult onset	AR	<i>CYP11B1</i>	99	1/50	1/5000	1/100000000	1/20000

Retinitis pigmentosa 2	XL	<i>RP2</i>	99	3/100	1/3333	1/44444444	1/13333
Retinitis pigmentosa 3	XL	<i>RPGR</i>	22	1/259	1/332	1/441032	1/1328
Retinitis pigmentosa-12	AR	<i>CRB1</i>	99	1/104	1/10400	1/432640000	1/41600
Retinitis pigmentosa 19	AR	<i>ABCA4</i>	99	3/100	1/3333	1/44444444	1/13333
Retinitis pigmentosa 20	AR	<i>RPE65</i>	84	1/228	1/1425	1/8122500	1/5700
Retinitis pigmentosa 25	AR	<i>EYS</i>	73	1/66	1/244	1/239012	1/978
Retinitis pigmentosa 26	AR	<i>CERKL</i>	97	1/148	1/4933	1/97351111	1/19733
Retinitis pigmentosa 28	AR	<i>FAM161A</i>	99	1/296	1/29600	1/3504640000	1/118400
Retinitis pigmentosa-40	AR	<i>PDE6B</i>	99	3/100	1/3333	1/44444444	1/13333
Retinitis pigmentosa 43	AR	<i>PDE6A</i>	99	3/100	1/3333	1/44444444	1/13333
Retinitis pigmentosa 47	AR	<i>SAG</i>	99	3/100	1/3333	1/44444444	1/13333
Retinitis pigmentosa 59	AR	<i>DHDDS</i>	99	1/296	1/29600	1/3504640000	1/118400
Microphthalmia with coloboma 3	AR	<i>VSX2</i>	99	1/91	1/9100	1/331240000	1/36400
Corneal endothelial dystrophy, autosomal recessive	AR	<i>SLC4A11</i>	99	<1/500	1/50000	1/10000000000	1/200000
Retinoschisis	XL	<i>RS1</i>	85	<1/500	1/3333	1/44444444	1/13333
Leber congenital amaurosis 13	AR/AD	<i>RDH12</i>	60	<1/500	1/1250	1/6250000	1/5000
Gyrate atrophy of choroid and retina with or without ornithinemia	AR	<i>OAT</i>	87	<1/500	1/3846	1/59171598	1/15385
Retinitis pigmentosa 37	AR/AD	<i>NR2E3</i>	99	1/209	1/20900	1/1747240000	1/83600
Leber congenital amaurosis 5	AR	<i>LCA5</i>	93	1/500	1/7143	1/204081633	1/28571
Auditory System							
Deafness, autosomal recessive 2	AR	<i>MYO7A</i>	99	1/206	1/20600	1/1697440000	1/82400
Deafness, autosomal recessive 1A	AR	<i>GJB2</i>	99	1/80	1/8000	1/256000000	1/32000
Deafness, autosomal recessive 4, with enlarged vestibular aqueduct	AR	<i>SLC26A4</i>	99	1/80	1/8000	1/256000000	1/32000
Deafness, autosomal recessive 77	AR	<i>LOXHD1</i>	99	1/500	1/50000	1/10000000000	1/200000
Deafness, autosomal recessive 1B	AR	<i>GJB6</i>	99	1/423	1/42300	1/7157160000	1/169200
Digestive System							
Cholestasis, progressive familial intrahepatic 2	AR	<i>ABCB11</i>	98	<1/500	1/25000	1/2500000000	1/100000
Cholestasis, progressive familial intrahepatic 3	AR/AD	<i>ABCB4</i>	99	<1/500	1/50000	1/10000000000	1/200000
Cholestasis, progressive familial intrahepatic 4	AR	<i>TJP2</i>	99	<1/502	1/50200	1/10080160000	1/200800
Cholestasis, progressive familial intrahepatic, 5	AR	<i>NR1H4</i>	99	<1/500	1/50000	1/10000000000	1/200000
Cholestasis, progressive familial intrahepatic 1	AR	<i>ATP8B1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Cystic fibrosis	AR	<i>CFTR</i>	95	<1/20	1/400	1/640000	1/1600
Liver failure, transient infantile	AR	<i>TRMU</i>	99	<1/500	1/50000	1/10000000000	1/200000
Diarrhea 1, secretory chloride, congenital	AR	<i>SLC26A3</i>	99	<1/500	1/50000	1/10000000000	1/200000
Hematologic System							
Factor XI deficiency, autosomal recessive	AR/AD	<i>F11</i>	99	1/500	1/50000	1/10000000000	1/200000
Hemophilia A	XL	<i>F8</i>	98	<1/500	1/25000	1/2500000000	1/100000
Hemophilia B	XL	<i>F9</i>	94	<1/500	1/8333	1/277777778	1/33333
Thalassemia, alpha-	AR	<i>HBA1/HBA2</i>	33/31	1/20	1/30; 1/29	1/3564; 1/3361	1/119; 1/116
Neutropenia, severe congenital 4, autosomal recessive	AR	<i>G6PC3</i>	99	1/341	1/34100	1/4651240000	1/136400
Fanconi anemia, complementation group A	AR	<i>FANCA</i>	93	1/239	1/3414	1/46629388	1/13657
Fanconi anemia, complementation group C	AR	<i>FANCC</i>	99	1/535	1/53500	1/11449000000	1/214000
Fanconi anemia, complementation group D2	AR	<i>FANCD2</i>	95	<1/200	1/4000	1/64000000	1/16000
Fanconi anemia, complementation group G	AR	<i>FANCG</i>	99	1/239	1/23900	1/2284840000	1/95600
Fanconi anemia, complementation group I	AR	<i>FANCI</i>	97	<1/200	1/6667	1/177777778	1/26667
Bernard-Soulier syndrome, type A1 (recessive)	AR	<i>GP1BA</i>	99	1/500	1/50000	1/10000000000	1/200000
Bernard-Soulier syndrome, type C	AR	<i>GP9</i>	99	1/500	1/50000	1/10000000000	1/200000
Thalassemia, beta	AR	<i>HBB</i>	99	1/158	1/15800	1/998560000	1/63200
Hypoprothrombinemia	AR	<i>F2</i>	99	1/33	1/3300	1/43560000	1/13200
Factor V deficiency	AR	<i>F5</i>	99	1/36	1/3600	1/51840000	1/14400
Abetalipoproteinemia	AR	<i>MTTP</i>	99	<1/500	1/50000	1/10000000000	1/200000

Thrombocytopenia, congenital amegakaryocytic	AR	<i>MPL</i>	99	1/102	1/10200	1/416160000	1/40800
Myopathy, lactic acidosis, and sideroblastic anemia 1	AR	<i>PUS1</i>	93	<1/500	1/7143	1/204081633	1/28571
Neutropenia, severe congenital 3, autosomal recessive	AR	<i>HAX1</i>	99	1/224	1/22400	1/2007040000	1/89600
Neutropenia, severe congenital, 5, autosomal recessive	AR	<i>VPS45</i>	99	1/224	1/22400	1/2007040000	1/89600
Multiple Systems							
Usher syndrome, type 1B	AR	<i>MYO7A</i>	99	1/206	1/20600	1/1697440000	1/82400
Usher syndrome, type 1C	AR	<i>USH1C</i>	99	1/353	1/35300	1/4984360000	1/141200
Usher syndrome, type 1D	AR	<i>CDH23</i>	99	1/285	1/28500	1/3249000000	1/114000
Usher syndrome, type 1F	AR	<i>PCDH15</i>	97	1/395	1/13167	1/693444444	1/52667
Usher syndrome, type 2A	AR	<i>USH2A</i>	99	1/126	1/12600	1/635040000	1/50400
Usher syndrome, type 2C	AR	<i>ADGRV1</i>	99	3/250	1/8333	1/277777778	1/33333
Usher syndrome, type 3A	AR	<i>CLRN1</i>	99	1/500	1/50000	1/10000000000	1/200000
Aicardi-Goutieres syndrome 5	AR	<i>SAMHD1</i>	95	<1/500	1/10000	1/400000000	1/40000
Arts syndrome	XL	<i>PRPS1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Bardet-Biedl syndrome 10	AR	<i>BBS10</i>	99	1/395	1/39500	1/6241000000	1/158000
Bardet-Biedl syndrome 12	AR	<i>BBS12</i>	99	1/791	1/79100	1/25027240000	1/316400
Bardet-Biedl syndrome 14	AR	<i>CEP290</i>	25	1/190	1/629	1/1584242	1/2517
Bardet-Biedl syndrome 1	AR	<i>BBS1</i>	99	1/367	1/36700	1/5387560000	1/146800
Bardet-Biedl syndrome 2	AR	<i>BBS2</i>	97	1/621	1/20700	1/1713960000	1/82800
Bardet-Biedl syndrome 3	AR	<i>ARL6</i>	93	<1/500	1/7143	1/204081633	1/28571
Bardet-Biedl syndrome 4	AR	<i>BBS4</i>	91	<1/500	1/5556	1/123456790	1/22222
Bardet-Biedl syndrome 6	AR	<i>MKKS</i>	99	<1/500	1/50000	1/10000000000	1/200000
Barth syndrome	XL	<i>TFAZZIN</i>	99	<3/1000	1/33333	1/4444444444	1/133333
Cohen syndrome	AR	<i>VPS13B</i>	96	<1/500	1/12500	1/625000000	1/50000
Ehlers-Danlos syndrome, kyphoscoliotic type, 1	AR	<i>PLOD1</i>	99	1/158	1/15800	1/998560000	1/63200
Ellis-van Creveld syndrome	AR	<i>EVC2</i>	98	1/240	1/12000	1/576000000	1/48000
Ellis-van Creveld syndrome	AR	<i>EVC</i>	93	1/142	1/2029	1/16460408	1/8114
Gitelman syndrome	AR	<i>SLC12A3</i>	99	1/100	1/10000	1/400000000	1/40000
Hermansky-Pudlak syndrome 1	AR	<i>HPS1</i>	91	<1/500	1/5556	1/123456790	1/22222
Hermansky-Pudlak syndrome 3	AR	<i>HPS3</i>	94	<1/500	1/8333	1/277777778	1/33333
Hermansky-Pudlak syndrome 4	AR	<i>HPS4</i>	99	<1/500	1/50000	1/10000000000	1/200000
Hermansky-Pudlak syndrome 5	AR	<i>HPS5</i>	99	<1/500	1/50000	1/10000000000	1/200000
Hermansky-Pudlak syndrome 6	AR	<i>HPS6</i>	85	<1/500	1/3333	1/444444444	1/13333
Joubert syndrome 17	AR	<i>CPLANE1</i>	81	1/472	1/2484	1/24685208	1/9937
Joubert syndrome 2	AR	<i>TMEM216</i>	99	1/472	1/47200	1/8911360000	1/188800
Joubert syndrome 3	AR	<i>AHI1</i>	57	1/472	1/1098	1/4819557	1/4391
Joubert syndrome 5	AR	<i>CEP290</i>	25	1/472	1/629	1/1584242	1/2517
Joubert syndrome 6	AR	<i>TMEM67</i>	70	1/472	1/1573	1/9901511	1/6293
Joubert syndrome 9	AR	<i>CC2D2A</i>	95	1/472	1/9440	1/356454400	1/37760
Laron dwarfism	AR	<i>GHR</i>	99	1/158	1/15800	1/998560000	1/63200
Meckel syndrome 1	AR	<i>MKS1</i>	99	1/260	1/26000	1/2704000000	1/104000
Netherton syndrome	AR	<i>SPINK5</i>	99	<2/100	1/5000	1/100000000	1/20000
Pendred syndrome	AR	<i>SLC26A4</i>	99	1/80	1/8000	1/256000000	1/32000
Shwachman-Diamond syndrome	AR	<i>SBDS</i>	99	1/146	1/14600	1/852640000	1/58400
Sjogren-Larsson syndrome	AR	<i>ALDH3A2</i>	99	1/250	1/25000	1/2500000000	1/100000
Smith-Lemli-Opitz syndrome	AR	<i>DHCR7</i>	99	1/30	1/3000	1/36000000	1/12000
Stuve-Wiedemann syndrome	AR	<i>LIFR</i>	80	<1/500	1/2500	1/25000000	1/10000
Usher syndrome, type 1G	AR	<i>USH1G</i>	99	1/600	1/60000	1/14400000000	1/240000
Wolcott-Rallison syndrome	AR	<i>EIF2AK3</i>	90	<1/500	1/5000	1/100000000	1/20000
Wolfram syndrome 1	AR	<i>WFS1</i>	99	<1/200	1/20000	1/1600000000	1/80000
Mental retardation-hypotonic facies syndrome, X-linked	XL	<i>ATRX</i>	58	<1/500	1/1190	1/5668934	1/4762

Bartter syndrome, type 4a	AR	<i>BSND</i>	99	1/500	1/50000	1/10000000000	1/200000
Mitochondrial complex III deficiency, nuclear type 1	AR	<i>BCS1L</i>	99	<1/500	1/50000	1/10000000000	1/200000
Bloom syndrome	AR	<i>BLM</i>	77	1/800	1/3478	1/48393195	1/13913
Multiple pterygium syndrome, lethal type	AR	<i>CHRNA</i>	99	<1/500	1/50000	1/10000000000	1/200000
Trichohepatoenteric syndrome 1	AR	<i>SKIC3</i>	99	1/500	1/50000	1/10000000000	1/200000
Osteoporosis-pseudoglioma syndrome	AR	<i>LRP5</i>	99	7/5000	1/71429	1/20408163265	1/285714
Mulibrey nanism	AR	<i>TRIM37</i>	80	1/100	1/500	1/1000000	1/2000
Cockayne syndrome, type A	AR	<i>ERCC8</i>	93	1/822	1/11743	1/551578776	1/46971
Cockayne syndrome, type B	AR	<i>ERCC6</i>	98	<1/500	1/25000	1/2500000000	1/100000
Roberts-SC phocomelia syndrome	AR	<i>ESCO2</i>	99	<1/500	1/50000	1/10000000000	1/200000
Joubert syndrome 7	AR	<i>RPGRIP1L</i>	87	1/259	1/1992	1/15877160	1/7969
Immunodeficiency-centromeric instability-facial anomalies syndrome 1	AR	<i>DNMT3B</i>	99	<1/500	1/50000	1/10000000000	1/200000
Nijmegen breakage syndrome	AR	<i>NBN</i>	86	1/158	1/1129	1/5094694	1/4514
Salla disease	AR	<i>SLC17A5</i>	95	<1/500	1/10000	1/400000000	1/40000
Ectodermal dysplasia 1, hypohidrotic, X-linked	XL	<i>EDA</i>	88	<1/500	1/4167	1/69444444	1/16667
Distal renal tubular acidosis 2 with progressive sensorineural hearing loss	AR	<i>ATP6V1B1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Alstrom syndrome	AR	<i>ALMS1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Steel syndrome	AR	<i>COL27A1</i>	99	<1/500	1/50000	1/10000000000	1/200000
Insensitivity to pain, congenital, with anhidrosis	AR	<i>NTRK1</i>	36	<1/500	1/781	1/2441406	1/3125
Mitochondrial DNA depletion syndrome 1 (MNGIE type)	AR	<i>TYMP</i>	99	<1/500	1/50000	1/10000000000	1/200000
Chediak-Higashi syndrome	AR	<i>LYST</i>	84	<1/500	1/3125	1/39062500	1/12500
Ehlers-Danlos syndrome, cardiac valvular type	AR	<i>COL1A2</i>	89	1/500	1/4545	1/82644628	1/18182
Schopf-Schulz-Passarge syndrome	AR	<i>WNT10A</i>	99	<1/500	1/50000	1/10000000000	1/200000
Joubert syndrome 4	AR	<i>NPHP1</i>	97	1/480	1/16000	1/1024000000	1/64000
Fragile X syndrome	XL	<i>FMR1</i>	99	1/250	1/25000	1/1250000000	1/33333

ACMG SF v3.2 list for reporting of secondary findings

Disorder	Inheritance	Gene/Locus	Category
Familial thoracic aortic aneurysm	AD	ACTA2	Cardiovascular
Hypertrophic cardiomyopathy	AD	ACTC1	Cardiovascular
Hereditary hemorrhagic telangiectasia	AD	ACVRL1	Miscellaneous
Familial adenomatous polyposis	AD	APC	Cancer
Familial hypercholesterolemia	AD	APOB	Cardiovascular
Wilson disease	AR	ATP7B	Miscellaneous
Dilated cardiomyopathy	AD	BAG3	Cardiovascular
Myofibrillar myopathy	AD	BAG3	Cardiovascular
Juvenile polyposis syndrome	AD	BMPR1A	Cancer
Hereditary breast and ovarian cancer	AD	BRCA1	Cancer
Hereditary breast and ovarian cancer	AD	BRCA2	Cancer
Biotinidase deficiency	AR	BTBD	Metabolic
Malignant hyperthermia	AD	CACNA1S	Miscellaneous
Long-QT syndrome type 14	AD	CALM1	Cardiovascular
Catecholaminergic polymorphic ventricular tachycardia	AD	CALM1	Cardiovascular
Long-QT syndrome type 15	AD	CALM2	Cardiovascular
Catecholaminergic polymorphic ventricular tachycardia	AD	CALM2	Cardiovascular
Long-QT syndrome type 16	AD	CALM3	Cardiovascular
Catecholaminergic polymorphic ventricular tachycardia	AD	CALM3	Cardiovascular
Catecholaminergic polymorphic ventricular tachycardia	AR	CASQ2	Cardiovascular
Ehlers-Danlos syndrome, vascular type	AD	COL3A1	Cardiovascular
Dilated cardiomyopathy	AD	DES	Cardiovascular
Myofibrillar myopathy	AD	DES	Cardiovascular
Arrhythmogenic right ventricular cardiomyopathy	AD	DSC2	Cardiovascular
Arrhythmogenic right ventricular cardiomyopathy	AD	DSG2	Cardiovascular
Arrhythmogenic right ventricular cardiomyopathy	AD	DSP	Cardiovascular
Dilated cardiomyopathy	AD	DSP	Cardiovascular
Hereditary hemorrhagic telangiectasia	AD	ENG	Miscellaneous
Marfan syndrome	AD	FBN1	Cardiovascular
Dilated cardiomyopathy	AD	FLNC	Cardiovascular
Hypertrophic cardiomyopathy	AD	FLNC	Cardiovascular
Myofibrillar myopathy	AD	FLNC	Cardiovascular
Pompe disease	AR	GAA	Metabolic
Fabry disease	XL	GLA	Cardiovascular Metabolic
Hereditary hemochromatosis (c.845G>A; p.C282Y homozygotes only)	AR	HFE	Miscellaneous
Maturity-Onset of Diabetes of the Young	AD	HNF1A	Miscellaneous
Long-QT syndrome type 2	AD	KCNH2	Cardiovascular
Long-QT syndrome type 1	AD	KCNQ1	Cardiovascular
Familial hypercholesterolemia	AD	LDLR	Cardiovascular
Dilated cardiomyopathy	AD	LMNA	Cardiovascular
Hereditary paraganglioma-pheochromocytoma syndrome	AD	MAX	Cancer
Multiple endocrine neoplasia type 1	AD	MEN1	Cancer
Lynch syndrome	AD	MLH1	Cancer
Lynch syndrome	AD	MSH2	Cancer
Lynch syndrome	AD	MSH6	Cancer
MUTYH-associated polyposis	AR	MUTYH	Cancer
Hypertrophic cardiomyopathy	AD	MYBPC3	Cardiovascular
Familial thoracic aortic aneurysm	AD	MYH11	Cardiovascular
Hypertrophic cardiomyopathy	AD	MYH7	Cardiovascular
Dilated cardiomyopathy	AD	MYH7	Cardiovascular
Hypertrophic cardiomyopathy	AD	MYL2	Cardiovascular
Hypertrophic cardiomyopathy	AD	MYL3	Cardiovascular
NF2-related schwannomatosis	AD	NF2	Cancer
Ornithine transcarbamylase deficiency	XL	OTC	Metabolic
Hereditary breast cancer	AD	PALB2	Cancer
Familial hypercholesterolemia	AD	PCSK9	Cardiovascular

Arrhythmogenic right ventricular cardiomyopathy	AD	<i>PKP2</i>	Cardiovascular
Lynch syndrome	AD	<i>PMS2</i>	Cancer
Hypertrophic cardiomyopathy	AD	<i>PRKAG2</i>	Cardiovascular Metabolic
<i>PTEN</i> hamartoma tumor syndrome	AD	<i>PTEN</i>	Cancer
Retinoblastoma	AD	<i>RB1</i>	Cancer
Dilated cardiomyopathy	AD	<i>RBM20</i>	Cardiovascular
Familial medullary thyroid cancer	AD	<i>RET</i>	Cancer
Multiple endocrine neoplasia type 2A	AD	<i>RET</i>	Cancer
Multiple endocrine neoplasia type 2B	AD	<i>RET</i>	Cancer
<i>RPE65</i> -related retinopathy	AR	<i>RPE65</i>	Miscellaneous
Malignant hyperthermia	AD	<i>RYR1</i>	Miscellaneous
Catecholaminergic polymorphic ventricular tachycardia	AD	<i>RYR2</i>	Cardiovascular
Long QT syndrome type 3	AD	<i>SCN5A</i>	Cardiovascular
Brugada syndrome	AD	<i>SCN5A</i>	Cardiovascular
Dilated cardiomyopathy	AD	<i>SCN5A</i>	Cardiovascular
Hereditary paraganglioma-pheochromocytoma syndrome	AD	<i>SDHAF2</i>	Cancer
Hereditary paraganglioma-pheochromocytoma syndrome	AD	<i>SDHB</i>	Cancer
Hereditary paraganglioma-pheochromocytoma syndrome	AD	<i>SDHC</i>	Cancer
Hereditary paraganglioma-pheochromocytoma syndrome	AD	<i>SDHD</i>	Cancer
Loeys-Dietz syndrome	AD	<i>SMAD3</i>	Cardiovascular
Juvenile polyposis syndrome	AD	<i>SMAD4</i>	Cancer
Hereditary hemorrhagic telangiectasia	AD	<i>SMAD4</i>	Miscellaneous
Peutz-Jeghers syndrome	AD	<i>STK11</i>	Cancer
Loeys-Dietz syndrome	AD	<i>TGFBR1</i>	Cardiovascular
Loeys-Dietz syndrome	AD	<i>TGFBR2</i>	Cardiovascular
Hereditary paraganglioma-pheochromocytoma syndrome	AD	<i>TMEM127</i>	Cancer
Arrhythmogenic right ventricular cardiomyopathy	AD	<i>TMEM43</i>	Cardiovascular
Dilated cardiomyopathy	AD	<i>TNNC1</i>	Cardiovascular
Hypertrophic cardiomyopathy	AD	<i>TNNI3</i>	Cardiovascular
Dilated cardiomyopathy	AD	<i>TNNT2</i>	Cardiovascular
Hypertrophic cardiomyopathy	AD	<i>TNNT2</i>	Cardiovascular
Li-Fraumeni syndrome	AD	<i>TP53</i>	Cancer
Hypertrophic cardiomyopathy	AD	<i>TPM1</i>	Cardiovascular
Catecholaminergic polymorphic ventricular tachycardia	AR	<i>TRDN</i>	Cardiovascular
Long QT syndrome	AR	<i>TRDN</i>	Cardiovascular
Tuberous sclerosis complex	AD	<i>TSC1</i>	Cancer
Tuberous sclerosis complex	AD	<i>TSC2</i>	Cancer
Dilated cardiomyopathy (truncating variants only)	AD	<i>TTN</i>	Cardiovascular
Hereditary transthyretin-related amyloidosis	AD	<i>TTR</i>	Miscellaneous
Von Hippel-Lindau syndrome	AD	<i>VHL</i>	Cancer
<i>WT1</i> -related Wilms tumor	AD	<i>WT1</i>	Cancer

Infertility panel

Disorder	Inheritance	Gene/Locus	Sex
Spermatogenic failure	<i>BRDT</i>	AR	M
Spermatogenic failure	<i>TSGA10</i>	AR	M
Spermatogenic failure	<i>FSIP2</i>	AR	M
Spermatogenic failure	<i>TTC21A</i>	AR	M
Spermatogenic failure	<i>DNAH1</i>	AR	M
Spermatogenic failure	<i>CFAP44</i>	AR	M
Spermatogenic failure	<i>SPATA16</i>	AR	M
Spermatogenic failure	<i>SPINK2</i>	AR	M
Spermatogenic failure	<i>SLC26A8</i>	AD	M
Spermatogenic failure	<i>ARMC2</i>	AR	M
Spermatogenic failure	<i>CFAP69</i>	AR	M
Spermatogenic failure	<i>TEX15</i>	AR	M
Spermatogenic failure	<i>NR5A1</i>	AD	M
Spermatogenic failure	<i>SOHLH1</i>	AD	M
Spermatogenic failure	<i>CFAP43</i>	AR	M
Spermatogenic failure	<i>NANOS1</i>	AD	M
Spermatogenic failure	<i>SYCE1</i>	AR	M
Spermatogenic failure	<i>CATSPER1</i>	AR	M
Spermatogenic failure	<i>PLCZ1</i>	AR	M
Spermatogenic failure	<i>DPY19L2</i>	AR	M
Spermatogenic failure	<i>SYCP3</i>	AD	M
Spermatogenic failure	<i>WDR66</i>	AR	M
Spermatogenic failure	<i>PPP2R3C</i>	AD	M
Spermatogenic failure	<i>FANCM</i>	AR	M
Spermatogenic failure	<i>AK7</i>	AR	M
Spermatogenic failure	<i>TDRD9</i>	AR	M
Spermatogenic failure	<i>MEIOB</i>	AR	M
Spermatogenic failure	<i>12-Sep</i>	AD	M
Spermatogenic failure	<i>PMFBP1</i>	AR	M
Spermatogenic failure	<i>ZMYND15</i>	AR	M
Spermatogenic failure	<i>KLHL10</i>	AD	M
Spermatogenic failure	<i>TEX14</i>	AR	M
Spermatogenic failure	<i>QRICH2</i>	AR	M
Spermatogenic failure	<i>TAF4B</i>	AR	M
Spermatogenic failure	<i>AURKC</i>	AR	M
Spermatogenic failure	<i>SUN5</i>	AR	M
Spermatogenic failure	<i>TEX11</i>	XLR	M
Spermatogenic failure	<i>USP9Y</i>	YL	M
Spermatogenic failure	<i>CEP19</i>	AR	M
Congenital bilateral absence of vas deferens, X-linked	<i>CFTR</i>	AR	M
Congenital bilateral absence of vas deferens, X-linked	<i>ADGRG2</i>	XL	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DRC1</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>ZMYND10</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAH1</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>CCDC39</i>	/	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAH5</i>	/	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>CCNO</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>RSPH9</i>	/	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>RSPH4A</i>	/	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>RSPH3</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAAF5</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAH11</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>NME8</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>SPAG1</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>LRRC6</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAI1</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>ARMC4</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>LRRC56</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAJB13</i>	AR	M

Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>CFAP300</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>CCDC65</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>KTU</i>	/	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAL1</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAAF4</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>HYDIN</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAAF1</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>GAS8</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAH9</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>GAS2L2</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>TTC25</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>CCDC103</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAI2</i>	/	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>CCDC40</i>	/	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>CCDC151</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>CCDC114</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>DNAAF3</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>C21orf59</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>RSPH1</i>	AR	M
Ciliary dyskinesia, primary, 1, with or without situs inversus	<i>PIH1D3</i>	XLR	M
Hypogonadotropic hypogonadism	<i>KISS1</i>	AR	M
Hypogonadotropic hypogonadism	<i>HS6ST1</i>	AD	M
Hypogonadotropic hypogonadism	<i>IL17RD</i>	AR/AD	M
Hypogonadotropic hypogonadism	<i>PROK2</i>	AD	M
Hypogonadotropic hypogonadism	<i>GNRHR</i>	AR	M
Hypogonadotropic hypogonadism	<i>TACR3</i>	AR	M
Hypogonadotropic hypogonadism	<i>SPRY4</i>	AD	M
Hypogonadotropic hypogonadism	<i>SEMA3A</i>	AD	M
Hypogonadotropic hypogonadism	<i>FEZF1</i>	AR	M
Hypogonadotropic hypogonadism	<i>FGF17</i>	AD	M
Hypogonadotropic hypogonadism	<i>GNRH1</i>	AR	M
Hypogonadotropic hypogonadism	<i>FGFR1</i>	AD	M
Hypogonadotropic hypogonadism	<i>CHD7</i>	AD	M
Hypogonadotropic hypogonadism	<i>NSMF</i>	AD	M
Hypogonadotropic hypogonadism	<i>FGF8</i>	AD	M
Hypogonadotropic hypogonadism	<i>WDR11</i>	AD	M
Hypogonadotropic hypogonadism	<i>FSHB</i>	AR	M
Hypogonadotropic hypogonadism	<i>TAC3</i>	AR	M
Hypogonadotropic hypogonadism	<i>DUSP6</i>	AD	M
Hypogonadotropic hypogonadism	<i>KISS1R</i>	AR	M
Hypogonadotropic hypogonadism	<i>LHB</i>	AR	M
Hypogonadotropic hypogonadism	<i>PROKR2</i>	AD	M
Hypogonadotropic hypogonadism	<i>FLRT3</i>	AD	M
Hypogonadotropic hypogonadism	<i>KAL1</i>	XLR	M
Hypogonadotropic hypogonadism	<i>RNF216</i>	AR	M
Hypogonadotropic hypogonadism	<i>SOX10</i>	AD	M
Hypogonadotropic hypogonadism	<i>SMCHD1</i>	AD	M
Hypogonadotropic hypogonadism	<i>NR0B1</i>	XLR	M
Hypogonadotropic hypogonadism	<i>DCAF17</i>	AR	M
Hypogonadotropic hypogonadism	<i>SLC29A3</i>	AR	M
Hypogonadotropic hypogonadism	<i>PROP1</i>	AR	M
Hypogonadotropic hypogonadism	<i>SOX2</i>	AD	M
Hypogonadotropic hypogonadism	<i>SEMA3E</i>	AD	M
Hypogonadotropic hypogonadism	<i>PNPLA6</i>	AR	M
Hypogonadotropic hypogonadism	<i>LEP</i>	AR	M
Hypogonadotropic hypogonadism	<i>LEPR</i>	AR	M
Hypogonadotropic hypogonadism	<i>LMNA</i>	AD	M
Hypogonadotropic hypogonadism	<i>POLR3A</i>	AR	M
Hypogonadotropic hypogonadism	<i>POLR3B</i>	AR	M
Androgen insensitivity	<i>AR</i>	XLR	M
Leydig cell hypoplasia	<i>LHCGR</i>	AR	M

Cryptorchidism	<i>INSL3</i>	AD	M
46XY sex reversal	<i>MAP3K1</i>	AD	M
46XY sex reversal	<i>ZFPM2</i>	AD	M
46XY sex reversal	<i>NR5A1</i>	AD	M
46XY sex reversal	<i>AKR1C2</i>	AR	M
46XY sex reversal	<i>AKR1C4</i>	AR	M
46XY sex reversal	<i>DHH</i>	AR	M
46XY sex reversal	<i>CBX2</i>	AR	M
46XY sex reversal	<i>NR0B1</i>	XL	M
46XY sex reversal	<i>SRY</i>	YL	M
46XY sex reversal	<i>SRD5A2</i>	AR	M
Persistent mullerian duct syndrome, type I and II	<i>AMHR2</i>	AR	M
Persistent mullerian duct syndrome, type I and II	<i>AMH</i>	AR	M
17-beta hydroxysteroid dehydrogenase III deficiency	<i>HSD17B3</i>	AR	M
Aarskog-Scott syndrome	<i>FGD1</i>	XLR	M
Prune belly syndrome	<i>CHRM3</i>	AR	M
Noonan syndrome	<i>NRAS</i>	AD	M
Noonan syndrome	<i>RIT1</i>	AD	M
Noonan syndrome	<i>PPP1CB</i>	AD	M
Noonan syndrome	<i>SOS1</i>	AD	M
Noonan syndrome	<i>RAF1</i>	AD	M
Noonan syndrome	<i>MRAS</i>	AD	M
Noonan syndrome	<i>BRAF</i>	AD	M
Noonan syndrome	<i>SHOC2</i>	AD	M
Noonan syndrome	<i>KRAS</i>	AD	M
Noonan syndrome	<i>PTPN11</i>	AD	M
Noonan syndrome	<i>SOS2</i>	AD	M
Noonan syndrome	<i>LZTR1</i>	AR	M
Hypogonadotropic hypogonadism	<i>KISS1</i>	AR	F
Hypogonadotropic hypogonadism	<i>HS6ST1</i>	AD	F
Hypogonadotropic hypogonadism	<i>IL17RD</i>	AR/AD	F
Hypogonadotropic hypogonadism	<i>PROK2</i>	AD	F
Hypogonadotropic hypogonadism	<i>GNRHR</i>	AR	F
Hypogonadotropic hypogonadism	<i>TACR3</i>	AR	F
Hypogonadotropic hypogonadism	<i>SPRY4</i>	AD	F
Hypogonadotropic hypogonadism	<i>SEMA3A</i>	AD	F
Hypogonadotropic hypogonadism	<i>FEZF1</i>	AR	F
Hypogonadotropic hypogonadism	<i>FGF17</i>	AD	F
Hypogonadotropic hypogonadism	<i>GNRH1</i>	AR	F
Hypogonadotropic hypogonadism	<i>FGFR1</i>	AD	F
Hypogonadotropic hypogonadism	<i>CHD7</i>	AD	F
Hypogonadotropic hypogonadism	<i>NSMF</i>	AD	F
Hypogonadotropic hypogonadism	<i>FGF8</i>	AD	F
Hypogonadotropic hypogonadism	<i>WDR11</i>	AD	F
Hypogonadotropic hypogonadism	<i>FSHB</i>	AR	F
Hypogonadotropic hypogonadism	<i>TAC3</i>	AR	F
Hypogonadotropic hypogonadism	<i>DUSP6</i>	AD	F
Hypogonadotropic hypogonadism	<i>KISS1R</i>	AR	F
Hypogonadotropic hypogonadism	<i>LHB</i>	AR	F
Hypogonadotropic hypogonadism	<i>PROKR2</i>	AD	F
Hypogonadotropic hypogonadism	<i>FLRT3</i>	AD	F
Hypogonadotropic hypogonadism	<i>KAL1</i>	XLR	F
Hypogonadotropic hypogonadism	<i>RNF216</i>	AR	F
Hypogonadotropic hypogonadism	<i>SOX10</i>	AD	F
Hypogonadotropic hypogonadism	<i>SMCHD1</i>	AD	F
Hypogonadotropic hypogonadism	<i>NR0B1</i>	XLR	F
Hypogonadotropic hypogonadism	<i>DCAF17</i>	AR	F
Hypogonadotropic hypogonadism	<i>SLC29A3</i>	AR	F
Hypogonadotropic hypogonadism	<i>PROP1</i>	AR	F
Hypogonadotropic hypogonadism	<i>SOX2</i>	AD	F
Hypogonadotropic hypogonadism	<i>SEMA3E</i>	AD	F

Hypogonadotropic hypogonadism	<i>PNPLA6</i>	AR	F
Hypogonadotropic hypogonadism	<i>LEP</i>	AR	F
Hypogonadotropic hypogonadism	<i>LEPR</i>	AR	F
Hypogonadotropic hypogonadism	<i>LMNA</i>	AD	F
Hypogonadotropic hypogonadism	<i>POLR3A</i>	AR	F
Hypogonadotropic hypogonadism	<i>POLR3B</i>	AR	F
Premature ovarian failure	<i>HFM1</i>	AR	F
Premature ovarian failure	<i>FIGLA</i>	AD	F
Premature ovarian failure	<i>FOXL2</i>	AD	F
Premature ovarian failure	<i>GDF9</i>	AR	F
Premature ovarian failure	<i>MSH5</i>	AR	F
Premature ovarian failure	<i>STAG3</i>	AR	F
Premature ovarian failure	<i>NOBOX</i>	AD	F
Premature ovarian failure	<i>NR5A1</i>	AD	F
Premature ovarian failure	<i>ERCC6</i>	AD	F
Premature ovarian failure	<i>SYCE1</i>	AR	F
Premature ovarian failure	<i>FANCM</i>	AR	F
Premature ovarian failure	<i>MCM8</i>	AR	F
Premature ovarian failure	<i>BMP15</i>	XL	F
Premature ovarian failure	<i>FLJ22792</i>	XLR	F
Premature ovarian failure	<i>DIAPH2</i>	XLD	F
Premature ovarian failure	<i>FMR1</i>	XL	F
Premature ovarian failure	<i>AARS2</i>	AR	F
Premature ovarian failure	<i>LMNA</i>	AD	F
Premature ovarian failure	<i>LARS2</i>	AR	F
Premature ovarian failure	<i>HSD17B4</i>	AR	F
Premature ovarian failure	<i>HARS2</i>	AR	F
Premature ovarian failure	<i>TWINK</i>	AR	F
Premature ovarian failure	<i>ERAL1</i>	AR	F
Premature ovarian failure	<i>CLPP</i>	AR	F
Premature ovarian failure	<i>AMH</i>	AR	F
Ovarian dysgenesis	<i>FSHR</i>	AR	F
Ovarian dysgenesis	<i>MRPS22</i>	AR	F
Ovarian dysgenesis	<i>MCM9</i>	AR	F
Ovarian dysgenesis	<i>SOHLH1</i>	AR	F
Ovarian dysgenesis	<i>NUP107</i>	AR	F
Ovarian dysgenesis	<i>ESR2</i>	AD	F
Ovarian dysgenesis	<i>PSMC3IP</i>	AR	F
Ovarian dysgenesis	<i>BMP15</i>	XL	F
Ovarian dysgenesis	<i>SAMD9</i>	AD	F
Leukoencephalopathy with vanishing white matter	<i>EIF2B4</i>	AR	F
Leukoencephalopathy with vanishing white matter	<i>EIF2B5</i>	AR	F
Leukoencephalopathy with vanishing white matter	<i>EIF2B2</i>	AR	F
Ovarian hyperstimulation syndrome	<i>FSHR</i>	AD	F
Oocyte maturation defect	<i>ZP3</i>	AD	F
Oocyte maturation defect	<i>WEE2</i>	AR	F
Oocyte maturation defect	<i>TUBB8</i>	AR/AD	F
Oocyte maturation defect	<i>ZP1</i>	AR	F
Oocyte maturation defect	<i>PANX1</i>	AD	F
Oocyte maturation defect	<i>PATL2</i>	AR	F
Oocyte maturation defect	<i>ZP2</i>	AR	F
Oocyte maturation defect	<i>ZP4</i>	/	F
Hydatidiform mole, recurrent	<i>KHDC3L</i>	AR	F
Hydatidiform mole, recurrent	<i>C11orf80</i>	/	F
Hydatidiform mole, recurrent	<i>NALP7</i>	AR	F
Hydatidiform mole, recurrent	<i>MEI1</i>	AR	F
Hydatidiform mole, recurrent	<i>TLE6</i>	AR	F
Hydatidiform mole, recurrent	<i>PADI6</i>	AR	F
Pregnancy loss, recurrent	<i>F5</i>	AD	F
Pregnancy loss, recurrent	<i>SYCP3</i>	AD	F
Pregnancy loss, recurrent	<i>ANXA5</i>	AD	F

Pregnancy loss, recurrent	<i>F2</i>	AD	F
Luteinizing hormone resistance	<i>LHCGR</i>	AR	F
46XY sex reversal	<i>MAP3K1</i>	AD	F
46XY sex reversal	<i>ZFPM2</i>	AD	F
46XY sex reversal	<i>NR5A1</i>	AD	F
46XY sex reversal	<i>AKR1C2</i>	AR	F
46XY sex reversal	<i>AKR1C4</i>	AR	F
46XY sex reversal	<i>DHH</i>	AR	F
46XY sex reversal	<i>CBX2</i>	AR	F
46XY sex reversal	<i>NR0B1</i>	XL	F
46XY sex reversal	<i>SRY</i>	YL	F
46XY sex reversal	<i>SRD5A2</i>	AR	F
Androgen insensitivity syndrome	<i>AR</i>	XLR	F
Cystic fibrosis	<i>CFTR</i>	AR	F
Polycystic ovary syndrome	<i>HSD17B3</i>	AR	F
Polycystic ovary syndrome	<i>INSR</i>	/	F
Polycystic ovary syndrome	<i>CYP11A1</i>	/	F
Polycystic ovary syndrome	<i>LHB</i>	AR	F
Polycystic ovary syndrome	<i>AR</i>	XLR	F
Polycystic ovary syndrome	<i>SRD5A2</i>	AR	F
Polycystic ovary syndrome	<i>CAPN10</i>	/	F
Polycystic ovary syndrome	<i>TH</i>	AR	F
Polycystic ovary syndrome	<i>PPARG</i>	AD	F
Polycystic ovary syndrome	<i>GDF9</i>	AR	F
Polycystic ovary syndrome	<i>AMH</i>	AR	F
Polycystic ovary syndrome	<i>HSD11B1</i>	AD	F
Polycystic ovary syndrome	<i>FSHR</i>	AR	F
Polycystic ovary syndrome	<i>LHCGR</i>	AR	F
Polycystic ovary syndrome	<i>LHB</i>	AR	F
Werner syndrome	<i>RECQL2</i>	AR	F
Adrenal hyperplasia, congenital, due to 21-hydroxylase deficiency	<i>CYP21A2</i>	AR	F
Hyperprolactinemia	<i>PRLR</i>	AR/AD	F