



Mackenzie's Mission Gene & Condition List

What conditions are being screened for in Mackenzie's Mission?

Genetic carrier screening offered through this research study has been carefully developed. It is focused on providing people with information about their chance of having children with a severe genetic condition occurring in childhood. The screening is designed to provide genetic information that is relevant and useful, and to minimise uncertain and unclear information.

How the conditions and genes are selected

The Mackenzie's Mission reproductive genetic carrier screen currently includes approximately 1300 genes which are associated with about 750 conditions. The reason there are fewer conditions than genes is that some genetic conditions can be caused by changes in more than one gene. The gene list is reviewed regularly.

To select the conditions and genes to be screened, a committee comprised of experts in genetics and screening was established including: clinical geneticists, genetic scientists, a genetic pathologist, genetic counsellors, an ethicist and a parent of a child with a genetic condition. The following criteria were developed and are used to select the genes to be included:

- Screening the gene is technically possible using currently available technology
- The gene is known to cause a genetic condition
- The condition affects people in childhood
- The condition has a serious impact on a person's quality of life and/or is life-limiting
 - For many of the conditions there is no treatment or the treatment is very burdensome for the child and their family. For some conditions very early diagnosis and treatment can make a difference for the child.

Types of conditions included

The conditions included in the screening vary in the way that they affect people and can involve one or many different parts of the body. Some of the ways that the conditions affect children can include:

Shortened life expectancy

Some conditions screened lead to a shortened life – either causing death in childhood, or with symptoms in childhood and early death in adulthood.

Intellectual disability

Some conditions cause intellectual disability which limits a person's ability to learn and develop independence. In some conditions this is severe – the child with the condition may never learn to walk or talk, whereas in others it is less severe – the child may be able to do many things for him or herself, but may need extra help and may not be able to live independently as an adult.

Physical conditions

Some conditions may affect the person physically, such as causing congenital heart disease or differences in how the limbs develop. In some cases these symptoms may be treatable, whereas in other cases there is no treatment available.

Neurological and muscular conditions

Some conditions are due to a problem with the brain itself, problems with the way the brain sends signals through the spinal cord and nerves to the body, or because the muscles themselves are weak. Sometimes these conditions can get worse over time.

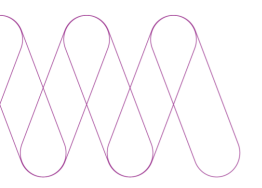
Important information about analysis and reporting of results

In addition to knowing what genes are being screened in Mackenzie's Mission, it is essential to also understand how the results are being analysed and reported. The screening is designed to be offered to a large number of people, with a focus on providing meaningful information that is useful to inform family planning.

Although a gene may be screened through Mackenzie's Mission, as outlined below, there are situations where particular genetic changes may not be analysed or reported:

A focus on severe conditions that occur in childhood

Some genetic conditions may vary in how much they affect people. This is because some genetic changes can have a more severe effect than others. Knowing about a chance of having a child with a mild form of a genetic condition often does not alter parents' reproductive plans and can cause confusion and distress. The focus of screening in this study is to provide information about the genetic chance of having a child with a severe



genetic condition. If a particular change in a gene is only associated with a mild form of the condition, this will not usually be reported to participants.

A 'couple screen'

In this study, a couple screening approach is used, meaning both biological parents of the pregnancy or planned pregnancy are screened at the same time. We are all genetic carriers for inherited conditions, however, many of the severe genetic conditions that occur in childhood are caused by **both** the biological mother and the biological father being carriers for the same autosomal recessive condition, or the biological mother being a carrier for an X-linked condition. Because of the very large number of genes screened, screening both biological parents at the same time and issuing a combined result provides the most useful information for that couple. If only one partner is a genetic carrier for an autosomal recessive condition/s, this will not be reported. This is because together, the couple will have a low chance of having a child with the condition. It is not practical to issue individual results for every person screened, and the results are most meaningful when combined together. If in the future either person has a new partner, that new couple should consider screening, as the results for the original couple are not relevant to the new couple.

A screening approach

There are many different types of genetic changes that can cause genetic conditions. It is important to understand that even with a 'low chance' result, there remains a small chance of a couple having a child with a genetic condition that was screened through Mackenzie's Mission. The genetic testing offered through this study is referred to as 'screening', because the technology used will detect many, but not all, genetic changes causing these conditions.

For fragile X syndrome and spinal muscular atrophy, targeted tests are used (each testing laboratory uses different methods which are described in the Mackenzie's Mission genetic carrier screening laboratory reports). In some circumstances, fragile X screening may also include AGG interruption analysis. For all other conditions, massively parallel sequencing is used. The testing techniques will not detect all genetic changes in each gene screened. For example, larger sections of extra or missing genetic material (called copy number variants, >50bp) or rearrangements will not be detected, which in some instances may be the main cause, or a major cause of a particular condition; examples include the *DMD*, *F8* and *TANGO2* genes. Additionally, in some cases this screening may not cover all genes associated with a particular genetic condition; this may be because the gene is associated with a mild form of the condition, or there are technical challenges in screening the gene.

Screening results are based on current knowledge

Knowledge about our genes is changing every day. Results from the genetic carrier screening performed through this study are being analysed and interpreted by experienced laboratory scientists. Their interpretation of the genetic information will be based on currently available information. So far, detailed genetic studies have not been done in people from all of the ethnic backgrounds found in the Australian population. This can make it more challenging to interpret some types of genetic results. For people from backgrounds for which there is less information, there may be a higher chance that couples who have an increased chance of having an affected child will not be identified.

When there is a family history of a genetic condition

While genetic carrier screening is relevant to everyone, regardless of whether there is a family history of a genetic condition, there will be some people who take part in this study who have a genetic condition themselves, or who have a relative/s with a genetic condition. It is important for people with a family history of a genetic condition who are wanting to have screening through Mackenzie's Mission to speak to a member of our study genetic counselling team, to determine whether the reproductive genetic carrier screen offered through this study is right for them. **Even if the gene causing the condition in their family is on the Mackenzie's Mission gene list, it is important to clarify whether the screening offered is able to detect the genetic change(s) present in that family.**

Please don't hesitate to contact our study team

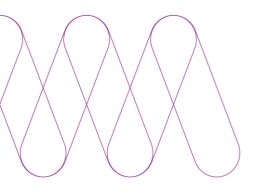
Our study team includes experienced genetic counsellors, clinical geneticists and laboratory scientists. We encourage healthcare providers and potential participants to contact us to discuss any queries they may have about the conditions screened through Mackenzie's Mission.

Mackenzie's Mission Study Team Email: info@mackenziesmission.org.au Phone: 1800 976 299 Website: mackenziesmission.org.au

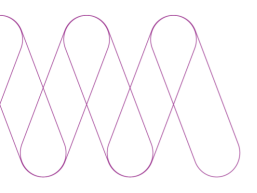
List of genes and conditions screened in Mackenzie's Mission

Please note that some genes appear on this list more than once, as changes in some genes can cause more than one different condition.

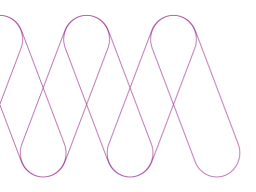
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Achalasia-addisonianism-alacrimia syndromeAAASAl Kaissi syndromeCDK10Athabaskan brainstem dysgenesis syndromeHOXA1Arthrogryposis, intellectual disability, and seizure disorderSLC35A33MC syndromeCOLEC11, MASP1Bardet-Biedl syndromeCOLEC11, MASP1Basel-Vanagait-Smirin-Yosef syndromeMED25Behr syndromeOPA1Boucher-Neuhauser syndromeHOXA1Brunner syndromeMAOAGoldberg-Shprintzen megacolon syndromeKIFBPBorjeson-Forssman-Lehmann syndromeBLMPartington syndromeARXPitt-Hopkins-like syndromeCNTNAP2Polyhydramnios, megalencephaly, and symptomatic epilepsySTRADAPERCHING syndromeKLHL7	Syndromes with intellectual	disability
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PERCHING syndrome KLHL7	Pitt-Hopkins-like syndrome	CNTNAP2
	Polyhydramnios, megalencephaly, and symptomatic epilepsy	STRADA
	PERCHING syndrome	KLHL7
Shaheen syndrome COG6	Shaheen syndrome	COG6
Growth retardation, intellectual developmental disorder, hypotonia, and hepatopathy	•	IARS1
Cataracts, growth hormone deficiency, sensory neuropathy, sensorineural hearing loss, and skeletal dysplasia (CAGSSS)		IARS2
Carey-Fineman-Ziter syndrome MYMK	Carey-Fineman-Ziter syndrome	МҮМК
Cerebellofaciodental syndrome BRF1	Cerebellofaciodental syndrome	BRF1
Craniofacial dysmorphism, skeletal anomalies, and intellectual disability syndrome TMCO1		TMCO1



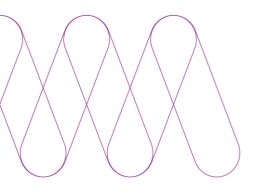
CHIME syndrome	PIGL
COACH syndrome	CC2D2A, RPGRIP1L, TMEM67
Cockayne syndrome	ERCC4, ERCC5, ERCC6, ERCC8
Cohen syndrome	VPS13B
Cerebrooculofacioskeletal syndrome (COFS)	ERCC2, ERCC6
Coffin-Lowry syndrome	RPS6KA3
Cowchock syndrome	AIFM1
De Sanctis-Cacchione syndrome	ERCC6
Developmental delay with short stature, dysmorphic features, and sparse hair	DPH1
Donnai-Barrow syndrome	LRP2
DOOR syndrome	TBC1D24
XFE progeroid syndrome	ERCC4
Desmosterolosis	DHCR24
Dyggve-Melchior-Clausen disease	DYM
Elsahy-Waters syndrome	CDH11
Fragile X syndrome	FMR1
Frontometaphyseal dysplasia	FLNA
Galloway-Mowat syndrome	WDR73, OSGEP
Gillespie syndrome	ITPR1
Griscelli syndrome	RAB27A
HSAN2D syndrome	SCN9A
Hypoparathyroidism-retardation-dysmorphism syndrome	TBCE
Hypotonia, infantile, with psychomotor retardation and characteristic facies	TBCK, UNC80, NALCN
Jawad syndrome	RBBP8
Jensen syndrome	TIMM8A
Johanson-Blizzard syndrome	UBR1
IFAP syndrome with or without BRESHECK syndrome	MBTPS2
Immunoskeletal dysplasia with neurodevelopmental abnormalities	EXTL3
Infantile liver failure syndrome	LARS1
Intellectual developmental disorder with dysmorphic facies, seizures, and distal limb anomalies	OTUD6B
Intellectual developmental disorder with cardiac arrhythmia	GNB5
Lujan-Fryns syndrome	MED12
Kohlschutter-tonz syndrome	ROGDI
Ohdo syndrome	MED12



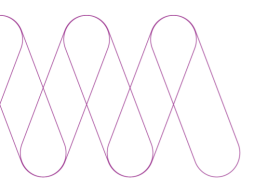
Opitz-Kaveggia syndrome	MED12
Opitz GBBB syndrome	MID1
Oliver-McFarlane syndrome	PNPLA6
Mosaic variegated aneuploidy syndrome	BUB1B
MEHMO syndrome	EIF2S3
Muscular dystrophy, congenital, with cataracts and intellectual disability	INPP5K
Nijmegen breakage syndrome	NBN, RAD50
Nance-Horan syndrome	NHS
Neurodevelopmental disorder with brain anomalies and additional features	PLAA, PRUNE1, VARS1, WDR45B
Multiple congenital anomalies-hypotonia-seizures syndrome	PIGA, PIGN, PIGT
Renpenning syndrome	PQBP1
Salt and pepper developmental regression syndrome	ST3GAL5
Seckel syndrome	ATR, CENPJ, CEP152, RBBP8
SESAME syndrome	KCNJ10
Smith-Lemli-Opitz syndrome	DHCR7
Spastic paraplegia and psychomotor retardation with or without seizures	HACE1
LIG4 syndrome	LIG4
Wieacker-Wolff syndrome	ZC4H2
Alacrima, achalasia, and intellectual disability syndrome	GMPPA
Chudley-McCullough syndrome	GPSM2
Growth retardation, developmental delay, coarse facies, and early death	FTO
Martsolf syndrome	RAB3GAP2
Pierson syndrome	LAMB2
Hemorrhagic destruction of the brain with subependymal calcification and cataracts	JAM3
Hennekam lymphangiectasia-lymphedema syndrome	CCBE1, FAT4
Perlman syndrome	DIS3L2
Temtamy preaxial brachydactyly syndrome	CHSY1
Filippi syndrome	CKAP2L
Fraser syndrome	FRAS1, FREM2
Orofaciodigital syndrome	CPLANE1, C2CD3, DDX59, SERPINH1, TMEM107, TCTN3
Roberts syndrome	ESCO2



SC phocomelia syndrome	ESCO2	
Warburg micro syndrome	RAB18, RAB3GAP1, RAB3GAP2	
Woodhouse-Sakati syndrome	DCAF17	
Van Maldergem syndrome	DCHS1, FAT4	
Warsaw breakage syndrome	DDX11	
You-Hoover-Fong syndrome	TELO2	
Syndromic microceph	aly	
Microcephaly, epilepsy, and diabetes syndrome	IER3IP1	
Microcephaly, progressive, seizures, and cerebral and cerebellar atrophy	QARS1	
Microcephaly-capillary malformation syndrome	STAMBP	
Microcephaly, short stature, and impaired glucose metabolism	TRMT10A	
Microcephaly, short-stature and endocrine dysfunction	XRCC4	
Microcephaly, short stature, and limb abnormalities	DONSON	
Microcephaly and chorioretinopathy	TUBGCP4, TUBGCP6	
Microcephaly, seizures, spasticity, and brain calcification	PCDH12	
X-linked syndromic intellectual disability		
Turner type	HUWE1	
Claes-Jensen type	KDM5C	
Christianson type	SLC9A6	
Siderius type	PHF8	
Type 35	RPL10	
Type 14	UPF3B	
CK syndrome	NSDHL	
Snyder-Robinson type	SMS	
Nascimento type	UBE2A	
Raymond type	ZDHHC9	
Intellectual disability, truncal obesity, retinal dystrophy, and micropenis	INPP5E	
Intellectual disability, X-linked, with cerebellar hypoplasia and distinctive facial appearance	OPHN1	
Syndromic brain malformations		
MASA syndrome	L1CAM	
CRASH syndrome	L1CAM	
Agenesis of the corpus callosum with peripheral neuropathy (Andermann syndrome)	SLC12A6	



Acrocallosal syndrome	KIF7
Proud syndrome	ARX
Temtamy syndrome	C12orf57
Cerebroretinal microangiopathy with calcifications and cysts	CTC1
Vici syndrome	EPG5
Proliferative vasculopathy and hydraencephaly-hydrocephaly syndrome	FLVCR2
Syndromic skin conditions with inte	ellectual disability
Cerebral dysgenesis, neuropathy, ichthyosis, and palmoplantar keratoderma syndrome	SNAP29
Adams-Oliver syndrome	DOCK6, EOGT
Syndromic vision conditions with in	tellectual disability
Peter's plus syndrome	B3GLCT
Congenital cataracts, hearing loss, and neurodegeneration	SLC33A1
Knobloch syndrome	COL18A1
Lowe syndrome	OCRL
Kaufman oculocerebrofacial syndrome	UBE3B
Kahrizi syndrome	SRD5A3
Optic atrophy with or without ataxia, intellectual disability, and seizures	RTN4IP1
Norrie disease	NDP
Syndromic growth conditions with in	tellectual disability
Simpson-Golabi-Behmel syndrome	OFD1, GPC3
Severe, lethal, neonatal sy	ndromes
Meckel syndrome	CC2D2A, CEP290, MKS1, NPHP3, RPGRIP1L, TMEM216, TMEM231, TMEM67
Alkuraya-Kucinskas syndrome	KIAA1109
Bowen-Conradi syndrome	EMG1
Fetal akinesia deformation sequence	RAPSN
Lethal congenital contracture syndrome	CNTNAP1, ERBB3, GLE1, GLDN, PIP5K1C
Ventriculomegaly with cystic kidney disease	CRB2
Hydrolethalus syndrome	HYLS1, KIF7
TARP syndrome	RBM10
Sudden infant death with dysgenesis of the testes syndrome	TSPYL1
Rigidity and multifocal seizure syndrome, lethal neonatal	BRAT1

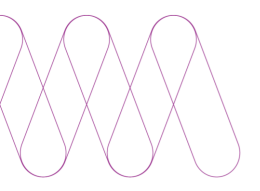


Syndromes without intellectual disability		
Multiple pterygium sync	Irome	
Lethal type	CHRNA1, RIPK4	
Escobar syndrome	CHRNG	
Multiple congenital abnor	malities	
Burn-McKeown syndrome	TXNL4A	
Bifid nose with or without anorectal and renal anomalies	FREM1	
Crisponi syndrome	CRLF1, CLCF1	
McKusick-Kaufman syndrome	MKKS	
Shwachman-Diamond syndrome	SBDS	
Split-hand foot malformation	WNT10B	
Werner syndrome	WRN	
VACTERL association X-linked	ZIC3	
Lipodystrophy, congenital generalized	BSCL2, CAVIN1	
Wolfram syndrome	CISD2, WFS1	
Urofacial syndrome	HPSE2, LRIG2	
Syndromic skin and skeletal conditions		
Rothmund-Thomson syndrome	RECQL4	
Alstrom syndrome	ALMS1	
GAPO syndrome	ANTXR1	
HELIX syndrome	CLDN10	
Haim-Munk syndrome	CTSC	
Laryngoonychocutaneous syndrome	LAMA3	
Miller syndrome	DHODH	
Macrocephaly, alopecia, cutis laxa, and scoliosis	RIN2	
Mandibuloacral dysplasia with type B lipodystrophy	ZMPSTE24	
Dyskeratosis congenita	DKC1, RTEL1, WRAP53	
Pigmentary disorder, reticulate, with systemic manifestations,	POLA1	
Papillon-Lefevre syndrome	CTSC	
Spondyloocular syndrome	XYLT2	
Treacher-Collins syndrome	POLR1C	
Schimke immunoosseous dysplasia	SMARCAL1	
Syndromic vision and bearing	anditiona	

Syndromic vision and hearing conditions

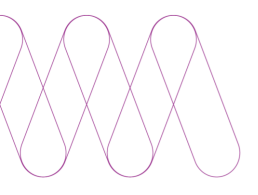
Usher syndrome

ADGRV1, CDH23, CIB2, CLRN1, MYO7A, PCDH15, USH1C, USH1G, USH2A, WHRN



Retinitis pigmentosa with skeletal anomalies	CWC27	
Jalili syndrome	CNNM4	
Syndromic vision and renal	conditions	
Senior-Loken syndrome	CEP290, NPHP1, NPHP4, SDCCAG8, IQCB1, WDR19	
Mitochondrial conditi	ons	
Conditions affecting multiple b	ody systems	
Combined oxidative phosphorylation deficiency	AARS2, C12orf65, CARS2, FARS2, ELAC2, GFM1, GTPBP3, MTFMT, MTO1, NARS2, RMND1, TSFM, TUFM, VARS2, TRIT1, EARS2	
Leigh and Leigh-like syn	drome	
Mitochondrial complex I deficiency	ACAD9, FOXRED1, NUBPL, NDUFA1, NDUFAF2, NDUFAF5, NDUFAF6, NDUFA10, NDUFA11, NDUFS6, NDUFS4, NDUFS2, NDUFS7, NDUFS8, NDUFS1, NDUFV1, NDUFV2	
Leigh syndrome due to cytochrome c oxidase deficiency	COX15	
Leigh syndrome, French Canadian type	LRPPRC	
Other mitochondrial conditions		
Mitochondrial complex II deficiency	SDHAF1	
Mitochondrial complex III deficiency	BCS1L, LYRM7, TTC19, UQCRC2, UQCRQ	
Mitochondrial complex IV deficiency	COX10, COA8, COX20, SURF1, PET100	
Mitochondrial complex V deficiency	TMEM70	
Mitochondrial DNA depletion syndrome	DGUOK, FBXL4, MGME1, MPV17, RRM2B, SUCLA2, SUCLG1, TK2, TWNK, TYMP	
Mitochondrial recessive ataxia syndrome (includes SANDO and SCAE)	TWNK	
Multiple mitochondrial dysfunctions syndrome	BOLA3, IBA57, ISCA2, NFU1	
Cardioencephalomyopathy, fatal infantile, due to cytochrome c oxidase deficiency 2	COX15, SCO2	
Sideroblastic anaemia with B-cell immunodeficiency, periodic fevers, and developmental delay	TRNT1	
Leukoencephalopathy with brain stem and spinal cord involvement and lactate elevation	DARS2	
Hyperuricemia, pulmonary hypertension, renal failure, and alkalosis (HUPRA syndrome)	SARS2	
HSD10 disease	HSD17B10	
Mohr-Tranebjaerg syndrome	TIMM8A	

Mitochondrial neurodevelopmental disorder, with abnormal movements and lactic acidosis	WARS2	
Myopathy, lactic acidosis, and sideroblastic anaemia	PUS1, LARS2, YARS2	
Mitochondrial short-chain enoyl-CoA hydratase 1 deficiency	ECHS1	
Lysosomal storage dis	orders	
Mannosidosis		
Alpha	MAN2B1	
Beta	MANBA	
Mucopolysaccharido	osis	
Mucopolysaccharidosis	GALNS, GNS, GUSB, IDS, IDUA	
Type VI (Maroteaux-Lamy)	ARSB	
Type IVB (Morquio)	GLB1	
Type IIIA (Sanfilippo A)	SGSH	
Type IIIB (Sanfilippo B)	NAGLU	
Type IIIC (Sanfilippo C)	HGSNAT	
Cystinosis		
Atypical nephropathic	CTNS	
Nephropathic	CTNS	
Late-onset juvenile or adolescent nephropathic	CTNS	
Ocular non-nephropathic	CTNS	
Other lysosomal storage disorders		
Galactosialidosis	CTSA	
Yunis-Varon syndrome	FIG4	
Fucosidosis	FUCA1	
Farber lipogranulomatosis	ASAH1	
Glycogen storage disease (Pompe)	GAA	
Geleophysic dysplasia	ADAMTSL2	
Krabbe disease	GALC, PSAP	
Fabry disease	GLA	
GM1-gangliosidosis	GLB1	
GM2-gangliosidosis	HEXA, GM2A	
Metachromatic leukodystrophy	ARSA, PSAP	
NA	GNPTAB, GNPTG, MCOLN1	
Mucolipidosis		
Polyglucosan body myopathy 1 with or without immunodeficiency	RBCK1	



Sandhoff disease	HEXB
Cirrhosis, cryptogenic	KRT8
Chediak-Higashi syndrome	LYST
Aspartylglucosaminuria	AGA
Schindler disease	NAGA
Sialidosis	NEU1
Combined SAP deficiency	PSAP
Marinesco-Sjogren syndrome	SIL1
Sialic acid storage disorder	SLC17A5
Niemann-Pick disease	NPC1, NPC2, SMPD1

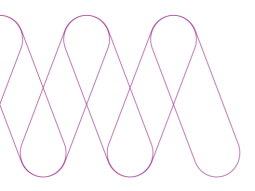
Metabolic conditions

Peroxisome biogenesis disorders

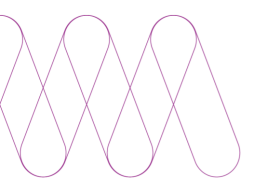
Including Zellweger syndrome, neonatal adrenoleukodystrophy
and infantile Refsum disease

PEX1, PEX10, PEX11B, PEX12, PEX13, PEX16, PEX2, PEX26, PEX3, PEX5, PEX6, PEX7

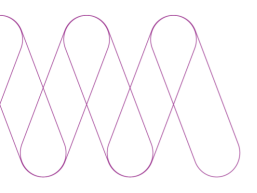
Organic acidemias		
Combined malonic and methylmalonic aciduria	ACSF3	
Argininosuccinic aciduria	ASL	
3-methylglutaconic aciduria	AUH, CLPB, DNAJC19, HTRA2, OPA3, SERAC1	
D-2-hydroxyglutaric aciduria	D2HGDH	
Glutaricaciduria	GCDH	
D-glyceric aciduria	GLYCTK	
L-2-hydroxyglutaric aciduria	L2HGDH	
Methylmalonic aciduria	MMADHC, MMUT	
Methylmalonic aciduria and homocystinuria	LMBRD1, MMACHC, MMADHC	
Alpha-methylacetoacetic aciduria	ACAT1	
Methylmalonic aciduria, vitamin B12-responsive	MMAA, MMAB	
Mevalonic aciduria	MVK	
Combined D-2- and L-2-hydroxyglutaric aciduria	SLC25A1	
Isovaleric acidemia	IVD	
Glutaric acidemia	ETFA, ETFB, ETFDH	
Other metabolic conditions		
Adenylosuccinase deficiency	ADSL	
Arts syndrome	PRPS1	
Chanarin-Dorfman syndrome	ABHD5	



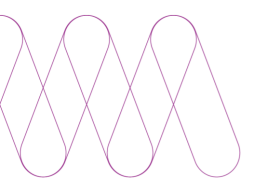
Glycogen storage disease	AGL, G6PC, GYS2, GBE1, LDHA, PFKM, SLC37A4
GABA-transaminase deficiency	ABAT
Fanconi-Bickel syndrome	SLC2A2
Hyperinsulinemic hypoglycemia	ABCC8, HADH, KCNJ11
Hyperoxaluria	AGXT
Hypermanganesemia with dystonia	SLC39A14
Succinic semialdehyde dehydrogenase deficiency	ALDH5A1
Fructose intolerance	ALDOB
Congenital disorders of glycosylation	ALG1, ALG11, ALG12, ALG3, ALG6, ALG8, ALG9, CCDC115, COG6, COG7, DOLK, DPAGT1, MGAT2, MPI, PGM1, PMM2, RFT1, SLC39A8, SSR4, SRD5A3, TMEM165
Congenital disorder of deglycosylation	NGLY1
Glycine encephalopathy	AMT, GLDC
Glycosylphosphatidylinositol biosynthesis defect	GPAA1
Argininemia	ARG1
Asparagine synthetase deficiency	ASNS
Canavan disease	ASPA
Citrullinemia	ASS1, SLC25A13
Chylomicron retention disease	SAR1B
Menkes disease and occipital horn syndrome	ATP7A
Maple syrup urine disease	BCKDHA, BCKDHB, DBT
Branched-chain ketoacid dehydrogenase kinase deficiency	BCKDK
GRACILE syndrome	BCS1L
Homocystinuria	CBS, MMADHC, MTHFR, MTR, MTRR
Lysinuric protein intolerance	SLC7A7
Proteinuria	CLCN5
Prolidase deficiency	PEPD
Hypomagnesemia	CLDN19, SLC30A10, TRPM6
Coenzyme Q10 deficiency	COQ2, COQ4, COQ6, COQ8A
Carbamoylphosphate synthetase I deficiency	CPS1
CPT 2 deficiency	CPT1A, CPT2
Methemoglobinemia	CYB5R3
Metabolic encephalomyopathic crises, recurrent, with rhabdomyolysis, cardiac arrhythmias, and neurodegeneration	TANGO2



Lipid storage myopathy due to flavin adenine dinucleotide synthetase deficiency	FLAD1
Peroxisomal acyl-CoA oxidase deficiency	ACOX1
17-alpha-hydroxylase deficiency	CYP17A1
17,20-lyase deficiency	CYP17A1
Cerebrotendinous xanthomatosis	CYP27A1
Aromatic L-amino acid decarboxylase deficiency	DDC
Dihydrolipoamide dehydrogenase deficiency	DLD
Wolcott-Rallison syndrome	EIF2AK3
Hypophosphatemic rickets	ENPP1
Hyperphosphatasia with intellectual disability syndrome	PIGV, PIGO, PGAP2, PGAP3
Ethylmalonic encephalopathy	ETHE1
Tyrosinemia	FAH, HPD
Fructose-1,6-bisphosphatase deficiency	FBP1
Fumarase deficiency	FH
Glutamate formiminotransferase deficiency	FTCD
Cerebral creatine deficiency syndrome	GAMT, GATM, SLC6A8
Gaucher disease	GBA, PSAP
Glycerol kinase deficiency	GK
Molybdenum cofactor deficiency	GPHN, MOCS1, MOCS2
Glutathione synthetase deficiency	GSS
3-hydroxyacyl-CoA dehydrogenase deficiency	HADH
LCHAD deficiency	HADHA
Trifunctional protein deficiency	HADHA, HADHB
Hemochromatosis	HAMP, HJV
Hemochromatosis 3-hydroxyisobutryl-CoA hydrolase deficiency	HAMP, HJV HIBCH
	· · ·
3-hydroxyisobutryl-CoA hydrolase deficiency	HIBCH
3-hydroxyisobutryl-CoA hydrolase deficiency Holocarboxylase synthetase deficiency	HIBCH HLCS
3-hydroxyisobutryl-CoA hydrolase deficiency Holocarboxylase synthetase deficiency HMG-CoA lyase deficiency	HIBCH HLCS HMGCL
 3-hydroxyisobutryl-CoA hydrolase deficiency Holocarboxylase synthetase deficiency HMG-CoA lyase deficiency HMG-CoA synthase-2 deficiency 	HIBCH HLCS HMGCL HMGCS2
 3-hydroxyisobutryl-CoA hydrolase deficiency Holocarboxylase synthetase deficiency HMG-CoA lyase deficiency HMG-CoA synthase-2 deficiency Lesch-Nyhan syndrome 	HIBCH HLCS HMGCL HMGCS2 HPRT1
 3-hydroxyisobutryl-CoA hydrolase deficiency Holocarboxylase synthetase deficiency HMG-CoA lyase deficiency HMG-CoA synthase-2 deficiency Lesch-Nyhan syndrome D-bifunctional protein deficiency 	HIBCH HLCS HMGCL HMGCS2 HPRT1 HSD17B4
 3-hydroxyisobutryl-CoA hydrolase deficiency Holocarboxylase synthetase deficiency HMG-CoA lyase deficiency HMG-CoA synthase-2 deficiency Lesch-Nyhan syndrome D-bifunctional protein deficiency Leprechaunism 	HIBCH HLCS HMGCL HMGCS2 HPRT1 HSD17B4 INSR
 3-hydroxyisobutryl-CoA hydrolase deficiency Holocarboxylase synthetase deficiency HMG-CoA lyase deficiency HMG-CoA synthase-2 deficiency Lesch-Nyhan syndrome D-bifunctional protein deficiency Leprechaunism Norum disease 	HIBCH HLCS HMGCL HMGCS2 HPRT1 HSD17B4 INSR LCAT



Cholesteryl ester storage disease	LIPA
Wolman disease	LIPA
Lipoyltransferase 1 deficiency	LIPT1
Lipoprotein lipase deficiency	LPL
Malonyl-CoA decarboxylase deficiency	MLYCD
Abetalipoproteinemia	MTTP
N-acetylglutamate synthase deficiency	NAGS
N-terminal acetyltransferase deficiency	NAA10
Ornithine transcarbamylase deficiency	OTC
Phenylketonuria (PKU)	PAH
Pyruvate carboxylase deficiency	PC
Hyperphenylalaninemia	PTS, QDPR, DNAJC12
Propionicacidemia	PCCA, PCCB
Proprotein convertase 1 deficiency	PCSK1
Pyruvate dehydrogenase deficiency	PDHA1, PDHB, PDP1
Phosphoglycerate kinase 1 deficiency	PGK1
Phosphoglycerate dehydrogenase deficiency	PHGDH
Refsum disease	PHYH
Pyruvate kinase deficiency	PKLR
Plasminogen deficiency	PLG
Dysplasminogenemia	PLG
Pyridoxamine 5'-phosphate oxidase deficiency	PNPO
Phosphoribosylpyrophosphate synthetase superactivity	PRPS1
Phosphoserine phosphatase deficiency	PSPH
Neu-Laxova syndrome	PHGDH, PSAT1
Riboflavin transport deficiency syndrome	SLC52A2, SLC52A3
Tumoral calcinosis, normophosphatemic	SAMD9
Lathosterolosis	SC5D
Emphysema-cirrhosis, due to AAT deficiency	SERPINA1
Hemorrhagic diathesis due to antithrombin Pittsburgh	SERPINA1
Monocarboxylate transporter 1 deficiency	SLC16A1
Thiamine metabolism dysfunction syndrome	SLC19A2, SLC19A3, SLC25A19, TPK1
Carnitine deficiency	SLC22A5
Hyperornithinemia-hyperammonemia-homocitrullinemia syndrome	SLC25A15
Acrodermatitis enteropathica	SLC39A4
Multiple sulfatase deficiency	SUMF1

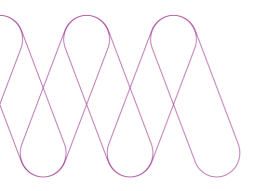


Salla disease	SLC17A5	
Sjogren-Larsson syndrome	ALDH3A2	
Sulfite oxidase deficiency	SUOX	
Transaldolase deficiency	TALDO1	
Barth syndrome	TAZ	
Adrenocorticotropic hormone deficiency	TBX19	
Transcobalamin II deficiency	TCN2	
Hemolytic anaemia due to triosephosphate isomerase deficiency	TPI1	
Crigler-Najjar syndrome	UGT1A1	
Orotic aciduria	UMPS	
Beta-ureidopropionase deficiency	UPB1	
VLCAD deficiency	ACADVL	
Wilson disease	ATP7B	
Endocrine conditions		

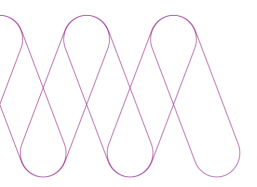
Congenital adrenal hyperplasia*	
Severe salt wasting type	CYP11A1, CYP11B2, NR0B1, POU1F1, PROP1, HSD3B2
Lipoid type	STAR

*Excludes 21-hydroxylase deficiency, as the CYP21A2 gene is not screened for technical reasons

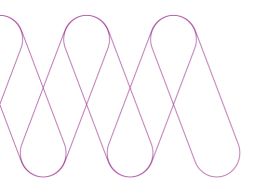
Diabetes mellitus	
Neonatal, with congenital hypothyroidism	GLIS3
Insulin-resistant, with acanthosis nigricans	INSR
Other endocrine condit	tions
Disordered steroidogenesis due to cytochrome P450 oxidoreductase	POR
Glucocorticoid deficiency	MC2R, MRAP, NNT
Growth hormone deficiency with pituitary anomalies	HESX1
Hyperparathyroidism, neonatal severe	CASR
Hypothryoidism, congenital	TSHB
Insulin-like growth factor resistance	IGF1R
Laron syndrome	GHR
Obesity, morbid, due to leptin deficiency	LEP
Pituitary hormone deficiency	HESX1, LHX3
Proopiomelanocortin (POMC) deficiency	POMC
Rabson-Mendenhall syndrome	INSR



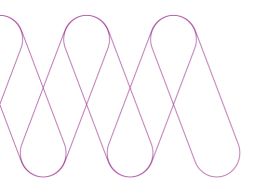
Neurological conditions		
White matter disorders		
Adrenoleukodystrophy	ABCD1	
Aicardi-Goutieres syndrome	ADAR, RNASEH2A, RNASEH2B, RNASEH2C, SAMHD1, TREX1	
Leukodystrophy, hypomyelinating	AIMP1, FAM126A, GJC2, HSPD1, POLR3A, POLR3B, PYCR2, RARS1, UFM1, VPS11	
Leukoencephalopathy with ataxia	CLCN2	
Leukoencephalopathy with vanishing white matter	EIF2B1, EIF2B2, EIF2B3, EIF2B4, EIF2B5	
Leukoencephalopathy, brain calcifications, and cysts	SNORD118	
Leukoencephalopathy, cystic, without megalencephaly	RNASET2	
Megalencephalic leukoencephalopathy with subcortical cysts	HEPACAM, MLC1	
Hypomyelination with brainstem and spinal cord involvement and leg spasticity (HBSL)	DARS1	
Pelizaeus-Merzbacher disease	PLP1	
Congenital brain malformations		
Pontocerebellar hypoplasia	AMPD2, CLP1, EXOSC3, EXOSC8, RARS2, SEPSECS, TBC1D23, TOE1, TSEN2, TSEN54, VPS53, VRK1	
Lissencephaly	ARX, KATNB1, LAMB1, NDE1, DCX, TMTC3	
Joubert syndrome	AHI1, ARL13B, CC2D2A, CEP290, CEP41, CPLANE1, CSPP1, INPP5E, KIF7, NPHP1, OFD1, RPGRIP1L, TCTN2, TCTN3, TMEM138, TMEM216, TMEM231, TMEM237, TMEM67	
Polymicrogyria	ADGRG1, RTTN, TUBA8	
Septooptic dysplasia	HESX1	
Band heterotopia	DCX, EML1	
Band-like calcification with simplified gyration and polymicrogyria	OCLN	
Cerebellar hypoplasia and intellectual disability with or without quadrupedal locomotion	VLDLR	
Periventricular heterotopia with microcephaly	ARFGEF2	
Poretti-Boltshauser syndrome	LAMA1	
Cortical malformations, occipital	LAMC3	
Infantile striatonigral degeneration	NUP62	



Microcephaly	
Isolated	ASPM, CDK5RAP2, CENPJ, CEP152, CIT, KIF14, KNL1, MCPH1, MFSD2A, MED17, PNKP, SLC25A19, STIL, WDR62, ZNF335
Hydrocephalus	
Non-syndromic hydrocephalus	L1CAM, CCDC88C, MPDZ
Hydrocephalus with congenital idiopathic intestinal pseudoobstruction	L1CAM
Hydrocephalus due to aqueductal stenosis	L1CAM
Hydrocephalus with Hirschsprung disease	L1CAM
Neurodegenerative conc	litions
Neuronal ceroid lipofuscinoses	CLN3, CLN5, CLN6, CLN8, CTSD, CTSF, MFSD8, PPT1, TPP1
Parkinson disease, juvenile-onset	DNAJC6, FBXO7, PLA2G6, ATP13A2
Encephalopathy, progressive	BSCL2, TBCD, NAXE
Moyamoya disease	GUCY1A1
Neurodegeneration with brain iron accumulation	C19orf12, PANK2, PLA2G6
Neurodegeneration due to cerebral folate transport deficiency	FOLR1
Neurodegeneration with ataxia, dystonia, and gaze palsy, childhood-onset	SQSTM1
PEHO syndrome	ZNHIT3
Infantile cerebellar-retinal degeneration	ACO2
Infantile neuroaxonal dystrophy 1	PLA2G6
Spastic tetraplegia, thin corpus callosum, and progressive microcephaly	SLC1A4
Troyer syndrome	SPART
Ataxias	
Ataxia-telangiectasia	ATM, MRE11
Ataxia, early-onset, with oculomotor apraxia and hypoalbuminemia	APTX
Ataxia, cerebellar, Cayman type	ATCAY
Ataxia, posterior column, with retinitis pigmentosa	FLVCR1
Ataxia-oculomotor apraxia 4	PNKP
Ataxia with isolated vitamin E deficiency	ТТРА
Cerebellar ataxia, cognitive disability, and disequilibrium (CAMRQ)	WDR81, ATP8A2
Spastic ataxia	KIF1C, MARS2, NKX6-2, SACS

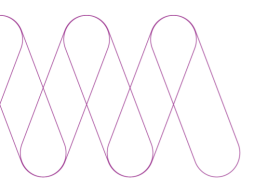


Spinocerebellar ataxia	GRM1, PMPCA, SETX, SNX14, STUB1, SCYL1, TPP1, WWOX	
Movement disorders	5	
Choreoacanthocytosis	VPS13A	
Dystonia	COL6A3	
Dystonia, dopa-responsive, due to sepiapterin reductase deficiency	SPR	
Dystonia, DOPA-responsive, with or without hyperphenylalaninemia	GCH1	
Parkinsonism-dystonia, infantile	SLC6A3	
Segawa syndrome	ТН	
Epilepsy		
Epilepsy, pyridoxine-dependent	ALDH7A1	
Epileptic encephalopathy, early infantile	AP3B2, ARV1, ARX, ARHGEF9, DENND5A, FRRS1L, MECP2, SLC13A5, SLC12A5, SLC25A22, TBC1D24, UBA5, WWOX	
Epilepsy, progressive myoclonic	CSTB, EPM2A, GOSR2, KCTD7, NHLRC1, PRICKLE1, SCARB2, TBC1D24	
Hyperekplexia	ATAD1, SLC6A5	
Epilepsy, early-onset, vitamin B6-dependent	PLPBP	
Epilepsy, X-linked, with variable learning disabilities and behaviour disorders	SYN1	
Epilepsy, hearing loss, and intellectual disability syndrome	SPATA5	
Cortical dysplasia-focal epilepsy syndrome	CNTNAP2	
Amish infantile epilepsy syndrome	ST3GAL5	
Intellectual disability	/	
Non-syndromic intellectual disability, X-linked	AP1S2, ARX, ATRX, BRWD3, CASK, CUL4B, CLCN4, DLG3, FTSJ1, GDI1, HCFC1, IL1RAPL1, IQSEC2, MECP2, NEXMIF, NLGN4X, PAK3, RAB39B, RLIM, SLC16A2, SYP, THOC2, TSPAN7, USP9X, ZNF711	
Non-syndromic intellectual disability, autosomal recessive	ADAT3, CC2D1A, ELP2, GPT2, HERC2, KPTN, LINS1, MAN1B1, MBOAT7, MED23, METTL23, NSUN2, PGAP1, PIGG, TRAPPC9, TTI2, TUSC3	
Other neurological conditions		
Sensorineural hearing loss, premature ovarian failure (females), variable intellectual disability, spasticity, ataxia	CLPP	

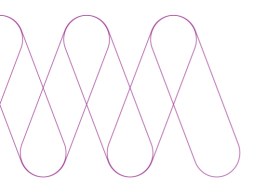


Cutaneous conditions		
Ichthyosis		
Ichthyosis, congenital, autosomal recessive	ABCA12, ALOX12B, ALOXE3, CERS3, CYP4F22, NIPAL4, TGM1	
Ichthyosis, leukocyte vacuoles, alopecia, and sclerosing cholangitis	CLDN1	
Epidermolytic hyperkeratosis	KRT10	
Cutis laxa		
Cutis laxa, autosomal recessive	ALDH18A1, ATP6V0A2, EFEMP2, FBLN5, LTBP4, PYCR1	
Ectodermal dysplasi	ia	
Ectodermal dysplasia, ectrodactyly and macular dystrophy	CDH3	
Ectodermal dysplasia	EDA, EDAR, IKBKG, KRT85	
Cutaneous conditions affecting the	nervous system	
Xeroderma pigmentosum	ERCC2, ERCC4, ERCC5, XPA, XPC	
Other cutaneous condit	tions	
Kindler syndrome	FERMT1	
Epidermolysis bullosa	COL7A1, COL17A1, DSP, ITGA6, ITGB4, KRT14, KRT5, LAMA3, LAMB3, LAMC2, PLEC	
Hyaline fibromatosis syndrome	ANTXR2	
Porokeratosis 3, disseminated superficial actinic	MVK	
Keratosis linearis with ichthyosis congenital and sclerosing keratoderma	POMP	
Netherton syndrome	SPINK5	
Poikilderma with neutropenia	USB1	
Restrictive dermopathy, lethal	LMNA, ZMPSTE24	
Triochthiodystrophy	ERCC2, GTF2H5, MPLKIP	
Transient bullous of the newborn	COL7A1	
Respiratory conditions		
Surfactant conditions		
Surfactant metabolism dysfunction, pulmonary	ABCA3, SFTPB	
Ciliary dyskinesia		
Ciliary dyskinesia, primary	ARMC4, CCDC103, CCDC114, CCDC39, CCDC40, CCNO, DNAAF1, DNAAF3, DNAAF4, DNAAF5, GAS8, HYDIN, LRRC6,	

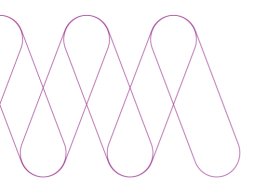
	PIH1D3, RSPH1, RSPH4A, RSPH9, SPAG1, ZMYND10
Ciliary dyskinesia, primary, with or without situs inversus	DNAH11, DNAH5, DNAI1, DNAI2
Other respiratory con	ditions
Cystic fibrosis	CFTR
Pulmonary veno-occlusive disease	EIF2AK4
Interstitial lung and liver disease	MARS1
Immunological cond	itions
Chronic granulomatous	disease
Deficiency of NCF-1	NCF1
Deficiency of NCF-2	NCF2
Deficiency of CYBA	СҮВА
X-linked	СҮВВ
Combined cellular and humoral immune defects with granulomas	RAG1, RAG2
Complement deficie	ncies
C1q	C1QA, C1QB, C1QC
C3	C3
C5	C5
C6	C6
C7	C7
C8	C8B
Factor D	CFD
Factor H	CFH
Factor I	CFI
Immunodeficienci	
Immunodeficiency	ATP6AP1, CARD11, CD3D, CD81, CTPS1, DOCK2, ICOS, IKBKB, IL12RB1, IL17RA, LRBA, MALT1, ORAI1, PGM3, RORC, SPNS1, STIM1, TNFRSF13B, TYK2
Mycobacteriosis	CYBB, IFNGR1, IFNGR2, STAT1
TCR-alpha/beta deficient	TRAC
Purine nucleoside phosphorylase deficiency	PNP
Hyper-IgM	CD40, CD40LG
Hyper-IgD syndrome	MVK
Hyper-IgE recurrent infection syndrome	DOCK8
Centromeric instability-facial anomalies syndrome	DNMT3B, ZBTB24



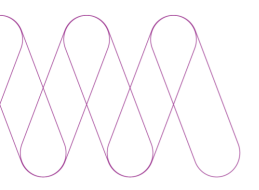
Combined immunodeficiency, moderate	IL2RG	
Combined immunodeficiency and megaloblastic anaemia with or without hyperhomocysteinemia	MTHFD1	
Immunoglobulin A deficiency	TNFRSF13B	
Neutropenia		
Severe, congenital	G6PC3, HAX1, JAGN1, VPS45, WAS	
Severe combined immunode	eficiencies	
Severe combined immunodeficiency	IL2RG	
Adenosine deaminase deficiency	ADA	
With microcephaly, growth retardation, and sensitivity to ionizing radiation	NHEJ1	
Athabascan type	DCLRE1C	
B cell-negative	RAG1, RAG2	
T-cell negative, B-cell/natural killer cell-positive type	IL7R, JAK3	
Reticular dysgenesis	AK2	
Other immunological conditions		
Agammaglobulinemia	BTK, IGHM	
Autoimmune disease, multisystem, with facial dysmorphism	ITCH	
Autoinflammation, lipodystrophy, and dermatosis syndrome	PSMB8	
Bone marrow failure syndrome	ERCC6L2, DNAJC21	
Bare lymphocyte syndrome	CIITA, RFXAP, TAP1	
Candidiasis, familial	CARD9	
Histiocytosis-lymphadenopathy plus syndrome	SLC29A3	
Hemophagocytic lymphohistiocytosis	PRF1, STX11, STXBP2, UNC13D	
Hepatic veno-occlusive disease with immunodeficiency	SP110	
Interleukin 1 receptor antagonist deficiency	IL1RN	
Immunodysregulation, polyendocrinopathy, and enteropathy	FOXP3	
Leukocyte adhesion deficiency	FERMT3, ITGB2	
Lymphoproliferative syndrome	CD27, ITK, SH2D1A, XIAP	
MHC class II deficiency, complementation group B	RFXANK	
Natural killer cell and glucocorticoid deficiency with DNA repair defect	MCM4	
Platelet abnormalities with eosinophilia and immune-mediated inflammatory disease	ARPC1B	
Properdin deficiency	CFP	
Pyogenic bacterial infections, recurrent, due to MYD88 deficiency	MYD88	



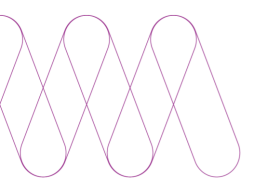
Selective T-cell defect	ZAP70	
T-cell immunodeficiency, congenital alopecia, and nail dystrophy	FOXN1	
Darsun syndrome	G6PC3	
Majeed syndrome	LPIN2	
Omenn syndrome	DCLRE1C, RAG1, RAG2	
Wiskott-Aldrich syndrome	WAS	
Gastrointestinal condit	ions	
Severe congenital diar	rhea	
With tufting enteropathy, congenital	EPCAM	
Secretory chloride, congenital	SLC26A3	
Secretory sodium, congenital,	SPINT2, SLC9A3	
Protein-losing enteropathy type	DGAT1	
Hepatic conditions	5	
Cholestasis, progressive familial intrahepatic	ABCB11, ABCB4, ATP8B1, TJP2	
Hepatic lipase deficiency	LIPC	
Porphyria	ALAD, UROS	
Liver failure, transient infantile	TRMU	
Hypercholanaemia	TJP2	
Other gastrointestinal conditions		
Microvillus inclusion disease	MYO5B	
Bile acid synthesis defect, congenital	AKR1D1, CYP7B1, HSD3B7	
Congenital short bowel syndrome	CLMP, FLNA	
Complement hyperactivation, angiopathic thrombosis, and protein-losing enteropathy	CD55	
Meconium ileus	GUCY2C	
Mitchell-Riley syndrome	RFX6	
Chronic atrial and intestinal dysrhythmia	SGO1	
Inflammatory bowel disease, congenital, severe	IL10RA	
Trichohepatoenteric syndrome	SKIV2L, TTC37	
Folate malabsorption, hereditary	SLC46A1	
Gastrointestinal defects and immunodeficiency syndrome	TTC7A	
Hyperbilirubinemia, familial transient neonatal	UGT1A1	
Haematological conditions		
Anaemia		
Sideroblastic, with ataxia	ABCB7	



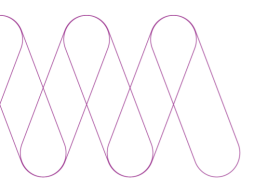
Anaemia, sideroblastic, pyridoxine-refractory	SLC25A38
Dyserythropoietic anaemia	SEC23B
Haemolytic anaemia due to hexokinase deficiency	HK1
Fanconi anaemia	ERCC4, FANCA, FANCB, FANCC, FANCD2, FANCE, FANCF, FANCG, FANCI, FANCL, UBE2T
Clotting conditions	5
Hypoprothrombinemia	F2
Factor V deficiency	F5
Factor VII deficiency	F7
Haemophilia A	F8
Haemophilia B	F9
Afibrinogenemia Dysfibrinogenemia Hypodysfibrinogenemia Hypofibrinogenemia	FGA, FGB, FGG
Combined factor V and VIII deficiency	LMAN1, MCFD2
Thrombotic thrombocytopenic purpura	ADAMTS13
Thrombocytopenia, congenital amegakaryocytic	MPL
Thrombophilia	PROC, PROS1
Thrombosis, hyperhomocysteinemic	CBS
von Willebrand disease	VWF
Thrombocytopenia, X-linked	WAS
Other haematological cor	ditions
Vitamin K-dependent clotting factors, combined deficiency of	VKORC1
Beta thalassemia	HBB
Sickle cell disease	HBB
Atransferrinemia	TF
Cardiovascular conditions	
Arrhythmias	
Ventricular tachycardia, catecholaminergic polymorphic	CASQ2
Jervell and Lange-Nielsen syndrome	KCNE1, KCNQ1
Ventricular tachycardia, catecholaminergic polymorphic with or without muscle weakness	TRDN



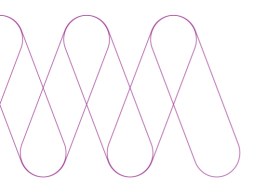
Cardiomyopathies	
Cardiomyopathy, dilated, with woolly hair and keratoderma (Naxos disease)	DSP, JUP
Dilated cardiomyopathy	FKTN
Structural cardiovascular c	onditions
Arterial calcification of infancy	ABCC6, ENPP1
Cardiac valvular dysplasia, X-linked	FLNA
Right atrial isomerism	GDF1
Hypoplastic left heart syndrome	GJA1
Arterial tortuosity syndrome	SLC2A10
Heterotaxy, visceral	ZIC3, MMP21
Congenital heart defects	ZIC3
Other cardiovascular conditions	
Sudden cardiac failure, infantile	PPA2
Renal conditions	
Syndromic renal conditions	
Alport syndrome	COL4A3, COL4A4, COL4A5
Dent disease	OCRL, CLCN5
Renal tubular acidosis with other abnormalities	ATP6V1B1, SLC4A4, SLC4A1
Bartter syndrome	BSND, CLCNKB, KCNJ1, SLC12A1
Renal-hepatic-pancreatic dysplasia	NPHP3, NEK8
Polycystic kidney and hepatic disease	PKHD1
Nephrotic syndrome	COQ8B, DGKE, LAMB2, NPHS1, NPHS2, NUP107, NUP93, PLCE1, SGPL1
Tubular conditions	
Renal tubular dysgenesis	ACE, AGT, REN
Renal tubular acidosis	ATP6V0A4
Other renal conditions	
Focal segmental glomerulosclerosis	CRB2
Pseudohypoaldosteronism	SCNN1A, SCNN1B
Nephronophthisis and related conditions	ANKS6, DCDC2, INVS, MAPKBP1, NPHP1, NPHP3, NPHP4, TMEM67, TTC21B, WDR19, XPNPEP3
Nephrogenic diabetes insipidus	AQP2



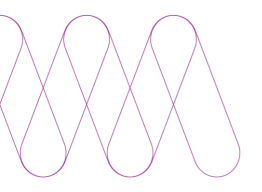
Neuromuscular conditions	
Atrophy	
Spinal muscular atrophy with progressive myoclonic epilepsy	ASAH1
Spinal muscular atrophy	SMN1, UBA1
Spinal muscular atrophy with congenital bone fractures	ASCC1
Arthrogryposis	
Arthrogryposis, distal	ECEL1, PIEZO2
Arthrogryposis lethal with anterior horn cell disease	GLE1
Arthrogryposis, renal dysfunction, and cholestasis	VIPAS39, VPS33B
Arthrogryposis multiplex congenita	LGI4
Dystrophy	
Limb-girdle muscular dystrophy	CAPN3, DYSF, PLEC, SGCA, SGCB, SGCD, SGCG, TCAP, TRAPPC11, TRIM32, TTN
Muscular dystrophy-dystroglycanopathy	B3GALNT2, CRPPA, FKRP, FKTN, GMPPB, LARGE1, POMGNT1, POMGNT2, POMK, POMT1, POMT2, RXYLT1
Muscular dystrophy, congenital	CHKB, LAMA2
Ullrich congenital muscular dystrophy	COL6A1, COL6A2, COL6A3
Duchenne muscular dystrophy	DMD *In VIC and NSW, most DMD carriers are unable to be detected due to limitations in testing technology
Becker muscular dystrophy	DMD *As above
Emery-Dreifuss muscular dystrophy	EMD, FHL1, LMNA
Muscular dystrophy, rigid spine	SELENON
Myopathy	
Myopathy, congenital	ACTA1
Nemaline myopathy	ACTA1, CFL2, KLHL40, KLHL41, LMOD3, NEB, TNNT1, TPM3
Myopathy, centronuclear, autosomal recessive	BIN1
Distal myopathy	DYSF
Myopathy with extrapyramidal signs	MICU1
Myopathy, X-linked	FHL1
Myopathy, X-linked, with excessive autophagy	VMA21
Inclusion body myopathy	GNE
Myopathy, areflexia, respiratory distress, and dysphagia, early- onset	MEGF10



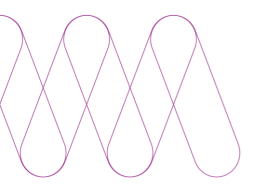
Myotubular myopathy, X-linked	MTM1
Minicore myopathy	RYR1
Myopathy, myofibrillar	KY, PYROXD1
Central core disease	RYR1
Myopathy, early-onset, with fatal cardiomyopathy	TTN
CAP myopathy	TPM3
Myasthenia	
Myasthenic syndrome	AGRN, ALG2, CHAT, CHRNA1, CHRND, CHRNE, COLQ, DOK7, DPAGT1, GFPT1, IGHMBP2, MUSK, RAPSN, SLC5A7
Neuropathy	
Charcot-Marie-Tooth disease	FGD4, FIG4, GDAP1, LMNA, LRSAM1, MFN2, MPZ, MTMR2, NDRG1, PRPS1, PRX, SBF2, SH3TC2
Dysautonomia, familial	ELP1
Insensitivity to pain, congenital	SCN9A, NTRK1
Neuromyotonia and axonal neuropathy	HINT1
Neuropathy, hereditary motor and sensory	HK1, IGHMBP2, KIF1A, SLC25A46
Neuropathy, hereditary sensory and autonomic	NGF, PRDM12, RETREG1, WNK1
Giant axonal neuropathy	GAN
Rhabdomyolysis	
Myoglobinuria, acute recurrent	LPIN1
Spasticity	
Spastic paralysis, infantile onset ascending	ALS2
Juvenile primary lateral sclerosis	ALS2
Spastic paraplegia	AP4M1, AP4B1, AP4S1, ATP13A2, ALDH18A1, B4GALNT1, CYP2U1, CYP7B1, DDHD2, DSTYK, FA2H, FARS2, GBA2, GJC2, KIF1A, NT5C2, PLP1, PNPLA6, SPG11, VPS37A, ZFYVE26
Connective tissue conditions	
Ehlers-Danlos syndrome (EDS)	
Ehlers-Danlos syndrome, progeroid type	ADAMTS2, B3GALT6, B4GALT7, PLOD1
Ehlers-Danlos syndrome, musculocontractural type	CHST14
Ehlers-Danlos syndrome with progressive kyphoscoliosis, myopathy, and hearing loss	FKBP14



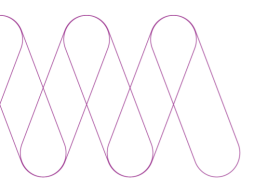
Vascular conditions	
Polyarteritis nodosa, childhood-onset	ADA2
Meester-Loeys syndrome	BGN
Ocular conditions	
Albinism	
Hermansky-Pudlak syndrome	HPS1, HPS3, HPS4, HPS5, HPS6
Oculocutaneous albinism	GPR143, LRMDA, OCA2, SLC24A5, SLC45A2, TYR, TYRP1
Dystrophies	
Retinal dystrophy, early-onset severe	LRAT, RCBTB1, CFAP410
Macular dystrophy with central cone involvement	MFSD8
Cone-rod dystrophy	AIPL1, ABCA4, C8orf37, CEP78, CNGB3, KCNV2, PDE6C, RPGRIP1, SEMA4A
Microphthalmia	
Isolated	ALDH1A3, RAX, VSX2
With coloboma	STRA6, VSX2
Syndromic	STRA6, RARB
Other ocular condition	ns
Achromatopsia	ATF6, CNGA3, CNGB3, GNAT2, OPN1LW
Aphakia	FOXE3
Congenital cataracts	AGK, FYCO1, NHS, TDRD7
Cone-rod synaptic disorder, congenital non-progressive	CABP4
Choroideremia	CHM
Congenital stationary night blindness	GPR179, NYX
Persistent hyperplastic primary vitreous	ATOH7
Macular degeneration (congenital)	CNGB3, RPGR
Leber congenital amaurosis	AIPL1, CEP290, CRB1, GUCY2D, LCA5, LRAT, NMNAT1, RD3, RDH12, RPE65, RPGRIP1, SPATA7, TULP1
Glaucoma (congenital)	CYP1B1
Peters anomaly	CYP1B1
Retinal arterial macroaneurysm with supravalvular pulmonic stenosis	IGFBP7
Retinitis pigmentosa	AGBL5, AIPL1, C8orf37, CRB1, DHDDS, FAM161A, IFT172, IMPG2, LRAT, MERTK, PDE6B, REEP6, RP2, SEMA4A, SPATA7, TULP1, USH2A



Progressive external ophthalmoplegia	POLG
Brittle cornea syndrome	PRDM5, ZNF469
Corneal opacification and other ocular anomalies	PXDN
Gaze palsy, horizontal, with progressive scoliosis	ROBO3
Foveal hypoplasia, with or without optic nerve misrouting and/or anterior segment dysgenesis	SLC38A8
Optic atrophy	TMEM126A
Skeletal conditions	5
Dysplasias	
Spondyloepiphyseal dysplasia with other abnormalities	CHST3, CCN6
Anauxetic dysplasia	POP1, RMRP
Spondyloepimetaphyseal dysplasia	B3GALT6, NANS
Desbuquois dysplasia	CANT1, XYLT1
Schneckenbecken dysplasia	SLC35D1
Short-rib thoracic dysplasia with or without polydactyly	CEP120, DYNC2LI1, KIAA0586, TTC21B, WDR34, WDR35, WDR60, DYNC2H1, IFT140, IFT172, IFT80, NEK1
Spondylometaepiphyseal dysplasia, short limb-hand type	DDR2
Spondylo-megaepiphyseal-metaphyseal dysplasia	NKX3-2
Chondrodysplasia, Grebe type	GDF5
Oculodentodigital dysplasia	GJA1
Smith-McCort dysplasia	DYM, RAB33B
Omodysplasia	GPC6
Dyssegmental dysplasia, Silverman-Handmaker type	HSPG2
Cranioectodermal dysplasia	IFT122
Opsismodysplasia	INPPL1
Otospondylomegaepiphyseal dysplasia	COL11A2, COL2A1
Greenberg skeletal dysplasia	LBR
Cleft lip/palate-ectodermal dysplasia syndrome	NECTIN1
Spondylometaphyseal dysplasia with additional abnormalities	PCYT1A, CFAP410
Chondrodysplasia, Blomstrand type	PTH1R
Metaphyseal dysplasia without hypotrichosis	RMRP
Craniolenticulosutural dysplasia	SEC23A
Langer mesomelic dysplasia	SHOX
De la Chapelle dysplasia	SLC26A2
Diastrophic dysplasia	SLC26A2



Chondrodysplasia punctata, rhizomelicAGPS, GNPAT, PEX7Mandibuloacral dysplasiaLMNAAcromesomelic dysplasiaHunter-Thompson typeGDF5Maroteaux typeNPR2Demirhan typeBMPR1BCCN6CranicosteoarthropathyHPGDHypertophic osteoarthropathyMMP2Cramptodact/tyl-arthropathy, and arthropathyMMP2Camptodact/tyl-arthropathy, craniofacial dysmorphism, and ongenital heart defectsBGGAT3Multicentic osteodysplasio, short stature, craniofacial dysmorphism, and ongenital heart defectsCCN7, RNU4ATACShort stature, onychodysplasia, facial dysmorphism, and hypotrichosisPCC1ANulticentic osteodysplasin, facial dysmorphism, and hypotrichosisPCC1AMultiper namismTRIM37Multiper namismCDther skeletal conditiumAntley-Bixler syndromePORHypoptrosphosphatasia, infantileALPLDiaphanospondylodysotosisBMPERMeler-Gorlin syndromeCA2, CLN7, OSTM1, TNERSF111Fibrochondrogenesis imperfecta, recessive typeCA2, CLN7, OSTM1, TNERSF11FibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCTAP, FKBP10, P3H1, SERPINF1, WNT1PypodysotosisDL13, HES7, MESP2Ellis-van Creveld syndromeFCK, CC/C2Raine syndromeFCK, CC/C2Raine syndromeFCK, CC/C2Raine syndromeFCK, CC/C2Raine syndromeFCK, CC/C2Raine syndromeFCK, CC/C2Raine syndromeFCK, CC/C2 <th>Craniofrontonasal dysplasia</th> <th>EFNB1</th>	Craniofrontonasal dysplasia	EFNB1	
Acromesomelic dysplasiaHunter-Thompson typeGDF5Maroteaux typeNPR2Demirhan typeBMPR1BArthropathiesArthropathy, progressive pseudorheumatoidCCN6CranicosteoarthropathyHPGDHypertrophic osteoarthropathyHPGDHypertrophic osteoarthropathyMMP2Camptodactyly-arthropathy-coxa vara-pericarditis syndromePRG4Multicentric osteolysis, nodulosis, and arthropathyB3GAT3Amelogenesis imperfecta and short stature, craniofacial dysmorphism, and oongenital heart defectsLTBP3Microcephalic osteodysplasia, facial dysmorphism, and hypotrichosisPCNT, RNU4ATACShort stature, onychodysplasia, facial dysmorphism, and hypotrichosisPCC1AShort stature, optic nerve atrophy, and Pelger-Huet anomalyNBASMultiper nanismTRIM37J-M syndromeCCDC8, OBSL1, CUL7Antley-Bixler syndromePORHypophosphatasia, infantileALPLDiaphanospondylodysotosisBMPERMeier-Gorlin syndromeCDT1, CDC45, ORC1, ORC6Osteopertosis, infantileCD111, CDC45, ORC1, ORC6Osteopertosis, infantileCTAP, FKBP10, P3H, SEPINF1, WNT1FibrochondrogenesisCTSKSpondylocostal dysotosisDLL3, HES7, MESP2Elis-van Creveld syndromeEVC, EVC2	Chondrodysplasia punctata, rhizomelic	AGPS, GNPAT, PEX7	
Hunter-Thompson typeGDF5Maroteaux typeNPR2Demirhan typeBMPR1BArthropathiesArthropathy, progressive pseudorheumatoidCCN6CranioosteoarthropathyHPGDHypertrophic osteoarthropathyHPGDHypertrophic osteoarthropathyMMP2Camptodactyly-arthropathy-coxa vara-pericarditis syndromePRG4Multicentric osteolysis, nodulosis, and arthropathyB3GAT3Annelogenesis imperfecta and short stature, craniofacial dysmorphism, and congenital heart defectsLTBP3Microcephalic osteodysplastic primordial dwarfismPCC1AShort stature, onychodysplasia, facial dysmorphism, and hypotrichosisPOC1AShort stature, onychodysplasia, facial dysmorphism, and hypotrichosisPOC1AMulticenter, optic nerve atrophy, and Pelger-Huet anomalyNBASMultipry nanismTRIM37Anteg-Bixler syndromeCCDC8, OBSL1, CUL7Antey-Bixler syndromePORHypophosphatasia, infatileALPLDiapanospondylodysotsoisBMPERMeier-Gorlin syndromeCD11, CDC45, ORC1, ORC6Osteopetrosis, infattileCA2, CLCN7, OSTM1, TCIRG1, TNFRSF11A, TNFSF11FibrochondrogenesisCTA1, NESF11A, TNFSF11FibrochondrogenesisCTA1, PCD45, ORC1, NRC61, TNFRSF11A, TNFSF11Shot stature, optic osteolysis, infantileCA2, CLCN7, OSTM1, TCIRG1, TNFRSF11A, TNFSF11FibrochondrogenesisCTA1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKB10, SH11, SERPINF1, WNT1PycnodysostosisCTSK	Mandibuloacral dysplasia	LMNA	
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Camptodactyly-arthropathy-coxa vara-pericarditis syndromePRG4Short stature and dwarfismB3GAT3Multiple joint dislocations, short stature, craniofacial dysmorphism, and congenital heart defectsB3GAT3Amelogenesis imperfecta and short statureLTBP3Microcephalic osteodysplastic primordial dwarfismPCNT, RNU4ATACShort stature, onychodysplasia, facial dysmorphism, and hypotrichosisPOC1AShort stature, optic nerve atrophy, and Pelger-Huet anomalyNBASMultiper nanismTRIM37Other skeletal conditions3-M syndromeCCDC8, OBSL1, CUL7Antley-Bixler syndromePORHypophosphatasia, infantileALPLDiaphanospondylodysostosisBMPRMeier-Gorlin syndromeCD11, CDC45, ORC1, ORC6Osteopetrosis, infantileCL11A1, COL11A2FibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Hypertrophic osteoarthropathy	HPGD	
Short stature and dwarfismMultiple joint dislocations, short stature, craniofacial dysmorphism, and congenital heart defectsB3GAT3Amelogenesis imperfecta and short statureLTBP3Microcephalic osteodysplastic primordial dwarfismPCNT, RNU4ATACShort stature, onychodysplasia, facial dysmorphism, and hypotrichosisPOC1AShort stature, optic nerve atrophy, and Pelger-Huet anomalyNBASMulibrey nanismTRIM37COther skeletal conditions3-M syndromeCCDC8, OBSL1, CUL7Antley-Bixler syndromePORHypophosphatasia, infantileALPLDiaphanospondylodysostosisBMPERMeier-Gorlin syndromeCDT1, CDC45, ORC1, ORC6Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Multicentric osteolysis, nodulosis, and arthropathy	MMP2	
Multiple joint dislocations, short stature, craniofacial dysmorphism, and congenital heart defectsB3GAT3Amelogenesis imperfecta and short statureLTBP3Microcephalic osteodysplastic primordial dwarfismPCNT, RNU4ATACShort stature, onychodysplasia, facial dysmorphism, and hypotrichosisPOC1AShort stature, optic nerve atrophy, and Pelger-Huet anomalyNBASMultibrey nanismTRIM37Other skeletal condition3-M syndromeCCDC8, OBSL1, CUL7Antley-Bixler syndromePORHypophosphatasia, infantileALPLDiaphanospondylodysostosisBMPERMeier-Gorlin syndromeCDT1, CDC45, ORC1, ORC6Osteopetrosis, infantileCL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Camptodactyly-arthropathy-coxa vara-pericarditis syndrome	PRG4	
and congenital heart defectsB3GAT3Amelogenesis imperfecta and short statureLTBP3Microcephalic osteodysplastic primordial dwarfismPCNT, RNU4ATACShort stature, onychodysplasia, facial dysmorphism, and hypotrichosisPOC1AShort stature, optic nerve atrophy, and Pelger-Huet anomalyNBASMulibrey nanismTRIM37CDCD28, OBSL1, CUL7Antley-Bixler syndromeCCDC8, OBSL1, CUL7Antley-Bixler syndromePORHypophosphatasia, infantileALPLDiaphanospondylodysostosisBMPERMeier-Gorlin syndromeCOL11, CDC45, ORC1, ORC6Osteopetrosis, infantileCOL11A1, COL11A2PibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Short stature and dwarf	fism	
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Short stature, onychodysplasia, facial dysmorphism, and hypotrichosisPOC1AShort stature, optic nerve atrophy, and Pelger-Huet anomalyNBASMulibrey nanismTRIM37Other skeletal conditions3-M syndromeCCDC8, OBSL1, CUL7Antley-Bixler syndromePORHypophosphatasia, infantileALPLDiaphanospondylodysostosisBMPERMeier-Gorlin syndromeCDT1, CDC45, ORC1, ORC6Osteopetrosis, infantileCA2, CLCN7, OSTM1, TCIRG1, TNFRSF11A, TNFSF11FibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Amelogenesis imperfecta and short stature	LTBP3	
hypotrichosisPOCTAShort stature, optic nerve atrophy, and Pelger-Huet anomalyNBASMulibrey nanismTRIM37Other skeletal conditions3-M syndromeCCDC8, OBSL1, CUL7Antley-Bixler syndromePORHypophosphatasia, infantileALPLDiaphanospondylodysostosisBMPERMeier-Gorlin syndromeCDT1, CDC45, ORC1, ORC6Osteopetrosis, infantileCA2, CLCN7, OSTM1, TCIRG1, TNFRSF11A, TNFSF11FibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Microcephalic osteodysplastic primordial dwarfism	PCNT, RNU4ATAC	
Mulibrey nanismTRIM37Other skeletal conditions3-M syndromeCCDC8, OBSL1, CUL7Antley-Bixler syndromePORHypophosphatasia, infantileALPLDiaphanospondylodysostosisBMPERMeier-Gorlin syndromeCDT1, CDC45, ORC1, ORC6Osteopetrosis, infantileCDT1, CDC45, ORC1, ORC6Osteopetrosis, infantileCOL11A1, COL11A2FibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2		POC1A	
Other skeletal conditions3-M syndromeCCDC8, OBSL1, CUL7Antley-Bixler syndromePORHypophosphatasia, infantileALPLDiaphanospondylodysostosisBMPERMeier-Gorlin syndromeCDT1, CDC45, ORC1, ORC6Osteopetrosis, infantileCA2, CLCN7, OSTM1, TCIRG1, TNFRSF11A, TNFSF11FibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Short stature, optic nerve atrophy, and Pelger-Huet anomaly	NBAS	
3-M syndromeCCDC8, OBSL1, CUL7Antley-Bixler syndromePORHypophosphatasia, infantileALPLDiaphanospondylodysostosisBMPERMeier-Gorlin syndromeCDT1, CDC45, ORC1, ORC6Osteopetrosis, infantileCA2, CLCN7, OSTM1, TCIRG1, TNFRSF11A, TNFSF11FibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Mulibrey nanism	TRIM37	
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DiaphanospondylodysostosisBMPERDiaphanospondylodysostosisCDT1, CDC45, ORC1, ORC6Meier-Gorlin syndromeCDT1, CDC45, ORC1, ORC6Osteopetrosis, infantileCA2, CLCN7, OSTM1, TCIRG1, TNFRSF11A, TNFSF11FibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Antley-Bixler syndrome	POR	
Meier-Gorlin syndromeCDT1, CDC45, ORC1, ORC6Osteopetrosis, infantileCA2, CLCN7, OSTM1, TCIRG1, TNFRSF11A, TNFSF11FibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Hypophosphatasia, infantile	ALPL	
Osteopetrosis, infantileCA2, CLCN7, OSTM1, TCIRG1, TNFRSF11A, TNFSF11FibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Diaphanospondylodysostosis	BMPER	
Osteopetrosis, infantileTNFRSF11A, TNFSF11FibrochondrogenesisCOL11A1, COL11A2Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Meier-Gorlin syndrome	CDT1, CDC45, ORC1, ORC6	
Osteogenesis imperfecta, recessive typeCRTAP, FKBP10, P3H1, SERPINF1, WNT1PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Osteopetrosis, infantile		
PycnodysostosisCTSKSpondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Fibrochondrogenesis	COL11A1, COL11A2	
Spondylocostal dysostosisDLL3, HES7, MESP2Ellis-van Creveld syndromeEVC, EVC2	Osteogenesis imperfecta, recessive type	CRTAP, FKBP10, P3H1, SERPINF1, WNT1	
Ellis-van Creveld syndrome EVC, EVC2	Pycnodysostosis	CTSK	
-	Spondylocostal dysostosis	DLL3, HES7, MESP2	
Raine syndrome FAM20C	Ellis-van Creveld syndrome	EVC, EVC2	
	Raine syndrome	FAM20C	



Bruck syndrome	FKBP10, PLOD2
Spondylocarpotarsal synostosis syndrome	FLNB
Brachydactyly	GDF5
Geroderma osteodysplasticum	GORAB
Craniosynostosis	IL11RA
Alazami syndrome	LARP7
Schwartz-Jampel syndrome	HSPG2
Stuve-Wiedemann syndrome/Schwartz-Jampel type 2 syndrome	LIFR
Acheiropody	LMBR1
Cenani-Lenz syndactyly syndrome	LRP4
Sclerosteosis	LRP4, SOST
Osteoporosis-pseudoglioma syndrome	LRP5
Orofacial cleft	NECTIN1
Brachyolmia 4 with mild epiphyseal and metaphyseal change	PAPSS2
Carpenter syndrome	RAB23, MEGF8
Baller-Gerold syndrome	RECQL4
RAPADILINO syndrome	RECQL4
Cartilage-hair hypoplasia	RMRP
Robinow syndrome	ROR2
Van den Ende-Gupta syndrome	SCARF2
Frank-ter Haar syndrome	SH3PXD2B
Achondrogenesis	SLC26A2, TRIP11
Atelosteogenesis	SLC26A2
Van Buchem disease	SOST
Kenny-Caffey syndrome	TBCE
Paget disease of bone	TNFRSF11B
Ulna and fibula, absence of, with severe limb deficiency	WNT7A
Fuhrmann syndrome	WNT7A
CODAS syndrome	LONP1
Keutel syndrome	MGP
Steel syndrome	COL27A1