

CGT 250 v2

Gene	Disease	Transcript	Mutations	Disease.description	products
ABCA4	Stargardt disease type 1; Cone-rod dystrophy type 3	NM_000350.2	NM_000350_2c.6449G>A, NM_000350_2c.6394G>T, NM_000350_2c.6320G>A, NM_000350_2c.6394G>T, NM_000350_2c.6320G>A, NM_000350_2c.61BC>T, NM_000350_2c.682G>A, NM_000350_2c.5912T>G, NM_000350_2c.582G>A, NM_000350_2c.5912T>G, NM_000350_2c.492F>T, NM_000350_2c.492F>T, NM_000350_2c.492F>T, NM_000350_2c.492F>T, NM_000350_2c.3210_3211dupCT, NM_000350_2c.3210_3211dupCT, NM_000350_2c.3210_3211dupCT, NM_000350_2c.3210_3211dupCT, NM_000350_2c.3916_3CT, NM_000350_2c.3916_3CT, NM_000350_2c.21616_3CT, NM_000350_2c.21616_3CT, NM_000350_2c.21616_3CT, NM_000350_2c.21616_3CT, NM_000350_2c.21616_3CT, NM_000350_2c.21616_3CT, NM_000350_2c.21616_3CT, NM_000350_2c.21616_3CT, NM_000350_2c.21616_3CT, NM_000350_2c.1616_3CT, NM_000350_2c.67363CT, NM_000350_2c.51016_3CT, NM_000350_2c.2636_3CT, NM_000350_2c.526_3CT, NM_000350_2c.536_3CT, NM_000350_2c.536_3CT, NM_000350_2c.536_3CT, NM_000350_2c.536_3CT, NM_000350_2c.536_3CT, NM_000350_2c.536_3CT, NM_	Stargardt disease type I follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ABCA4 gene located on chromosomal region Ip22. The age of onset is infantile. This disease is characterized by progressive central vision loss, mild loss of color vision, delayed dark adaptation and macular atrophy with or without paramacular flecks and degeneration of the underlying retinal pigment ephitelium. The estimated prevalence is 12,000-10,000. Mutations in the ABCA4 gene account also for 30 to 60 percent of cases of conerod dystrophy that are inherited in an autosomal recessive pattern. The problems associated with this condition include a loss of visual sharpness (acuity), an increased sensitivity to light (photophobia), and impaired color vision.	600,25
ACAD9	Mitochondrial complex I deficiency due to ACAD9	NM_014049.4	NM_014049.4:c.23deIT, NM_014049.4:c.130T>A, NM_014049.4:c.359deIT, NM_014049.4:c.453+IG>A, NM_014049.4:c.797G>A, NM_014049.4:c.976G>C, NM_014049.4:c.1240C>T, NM_014049.4:c.1249C>T, NM_014049.4:c.1594C>T	These vision problems worsen over time. Mitochondrial complex I deficiency due to ACAD9 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ACAD9 gene located on chromosomal region 3q213. The age of onset is neonatal/infantile. This disease is a multisystem disorder characterized by infantile onset of acute metabolic acidosis, hypertrophic cardiomyopathy, and muscle weakness associated with a deficiency of mitochondrial complex I activity in muscle, liver, and fibroblasts (summary by Haack et al., 2010). Medium-chain acyl-CoA dehydrogenase (MCAD) deficiency follows an	600,25
ACADM	Medium-chain acyl-CoA dehydrogenase deficiency	NM_001286043.1	NM_001286043.1:c.250C>T, NM_001286043.1:c.386-2A>G, NM_001286043.1:c.461C>T, NM_001286043.1:c.548_551delCTGA, NM_001286043.1:c.7546C>A, NM_001286043.1:c.715C>T, NM_001286043.1:c.756C>A, NM_001286043.1:c.835C>T, NM_001286043.1:c.956A>G, NM_001286043.1:c.898G>A, NM_001286043.1:c.916.928delCCAATCGGACTT, NM_001286043.1:c.1083delG, NM_001286043.1:c.1084A>G, NM_001286043.1:c.1201_1204delTTAG	autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ACADM gene located on chromosomal region 1p31. Inherited deficiency of MCAD is a condition that prevents the body from converting certain fats to energy, particularly during periods without food (fasting). Signs and symptoms of MCAD deficiency typically appear during infancy or early childhood and can include vomitting, lack of energy (lethargy), and low blood sugar (hypoglycemia). Individuals with MCAD deficiency are at risk of serious complications such as seizures, breathing difficulties, liver problems, brain damage, coma, and sudden death. The estimated prevalence is 14,990-127,000 in	
ACADS	Short-chain acyl-CoA dehydrogenase deficiency	NM_000017.3	NM_000017.3:c:136C>T, NM_000017.3:c:319C>T, NM_000017.3:c:417G>C, NM_000017.3:c:561_568delCAATGCCT, NM_000017.3:c:1095G>T, NM_000017.3:c:1147C>T	Caucasian populations and 114,600 in worldwide populations. Short-chain acyl-CoA dehydrogenase deficiency follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ACADS gene located on chromosomal region 12q24.31. The age of onset is infantile. This disease is characterized by seizures, developmental delay, failure to grow with poor feeding, and usually muscle weakness and hypotonia. The prevalence is <150,000. Short/branched-chain acyl-CoA dehydrogenase deficiency follows an autosomal recessive	
ACADSB	Short/branched-chain acyl- CoA dehydrogenase deficiency	NM_001609.3	NM_001609.3:c:303+1G>A, NM_001609.3:c:443C>T, NM_001609.3:c:621G>A, NM_001609.3:c:763C>T	pattern of inheritance and is caused by pathogenic variants in the ACADSB gene located on chromosomal region 10q26.3. The age of onset is neonatal/infantile. This disease is characterized by muscle hypotonia,	600,25

				cereprai paisy, developmental delay, lethargy, hypoglycemia, and metabolic acidosis. The prevalence is	
ACADVL	Very long-chain acyl-CoA dehydrogenase deficiency	NM_001270447.1	NM_001270447.1:c.347-1G>A, NM_001270447.1:c.367_368delCA, NM_0012704471:c.347-1G>A, NM_001270447.1:c.367_368delCA, NM_0012704471:c.362-3.	<11,000,000. <11,000,000. Very long-chain acyl-CoA dehydrogenase deficiency follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ACADVL gene located on chromosomal region 17p13.1. The age of onset is neonatal/infantile. This disease is characterized by cardiomyopathy, hypoketotic hypoglycemia, liver disease, exercise intolerance and rhabdomyolysis. The prevalence is 1:100,000-9:100,000. Renal tubular dysgenesis deficiency follows an autosomal recessive	600,25
ACE	Renal tubular dysgenesis	NM_000789.3	NM_000789.3:c:798C>G, NM_000789.3:c:1319_1322delTGGA, NM_000789.3:c:1486C>T, NM_000789.3:c:1511delC, NM_000789.3:c:1587 2A>G, NM_000789.3:c:2371C>T	pattern of inheritance and the most common cause are pathogenic variants in the ACE (chromosomal region 17q23.3). The age of onset is fetal. This disease is characterized by absent or poorly developed proximal tubules of the kidneys, persistent oligohydramios, leading	600,25
ADA	Adenosine deaminase deficiency / Severe combined immunodeficiency due to ADA deficiency	NM_000022.3	NM_000022.3:c:986C>T, NM_000022.3:c:956_960delAAGAG, NM_000022.3:c:890C>A, NM_000022.3:c:872C>T, NM_000022.3:c:320T>C	to Potter sequence, and skull ossification defects. Adenosine deaminase deficiency follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ADA gene located on chromosomal region 20q1312. The age of onset is neonatal/infantile. This disease is characterized by profound lymphopenia and very low immunoglobulin levels of all isotypes resulting in severe and recurrent opportunistic infections. The annual incidence is 1200,000-11,000,000. The	600,25
ADGRVI	Usher syndrome, type 2C	nm_032119.3	NM_032119.3:c.2258_2270delAAGTGCTGAAATC, NM_0321193:c.2864C>A, NM_0321193:c.5357_5358delAA, NM_0321193:c.6275-16-A, NM_0321193:c.6312dupt, NM_0321193:c.6901C>T, NM_0321193:c.6312dupt, NM_0321193:c.8790delC, NM_0321193:c.11377O>T, NM_0321193:c.14973- 1G>C, NM_0321193:c.15196.15199dupCAAA, NM_0321193:c.17668_17669delAT, NM_0321193:c.18131A>G	prevalence is 1:100,000-9:100,000. Usher syndrome type 2C follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ADGRVI and PDZD7 genes located on chromosomal regions 5q1.4.3 and 10q24.32 respectively. The age of onset is infantile. This disease is characterized by the association of sensorineural prelingual deafness (usually congenital) with retinitis pigmentosa and progressive vision loss. The	600,25
AGL	Clycogen storage disease type 3	NM_000028.2	NM_000028 2:c16C>T, NM_000028 2:c18.19delGA, NM_000028 2:c194-2A>T, NM_000028 2:c1222C>T, NM_000028 2:c1485delT, NM_000028 2:c1783C>T, NM_000028 2:c1999delC, NM_000028 2:c239G>A, NM_000028 2:c259G>T, NM_000028 2:c376-627delGA, NM_000028 2:c359G>A, NM_000028 2:c346-072A>G, NM_000028 2:c456G-15T, NM_000028 2:c4529dupA	prevalence is 1/30,000. Glycogen storage disease (CSD) type 3 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the AGL gene located on chromosomal region 1p21.2 The age of onset is infantile. This metabolic disorder is caused by deficiency of the glycogen debrancher enzyme and is associated with an accumulation of abnormal glycogen with short outer chains. Most patients are enzyme- deficient in both liver and muscle (IIIa), but about 15% are enzyme- deficient in liver only (IIIb) (Shen et al., 1996). These subtypes have been explained by differences in tissue expression of the deficient enzyme (Endo et al., 2006). In rare cases, selective loss of only 1 of the 2 debranching activities, glucosidase or transferase, results in type III or IIIId, respectively (Yan Hoof and Hers, 1967; Ding et al., 1990). Clinically, patients with GSD type 3 present in infancy or early childhood with hepatormegaly, hypoglycemia, and growth retardation. Muscle weakness in those with IIIs minimal in childhood but can become more severe in adults; some patients develop cardiomyopathy (Shen et al., 1996). Primary hyperoxaluria type I follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the AGXT gene	600,25

AGXT	Hyperoxaluria, primary, type 1	NM_000030.2	NM_000030.2:c.33dupC, NM_000030.2:c.121G>A, NM_000030.2:c.166-2A>G, NM_000030.2:c.246A>G, NM_000030.2:c.245A>G, NM_000030.2:c.3271>C, NM_000030.2:c.4547>A, NM_000030.2:c.456G>A, NM_000030.2:c.506C>A, NM_000030.2:c.506C>A, NM_000030.2:c.507 <a, nm_000030.2:c.507<a,="" nm_00030.2:c.507<a,="" nm_00030.2:c.731t="">C, NM_000030.2:c.738G>A</a,>	located on chromosomal region 2g37.3. The age of onset is variable. This disease is characterized by variable clinical presentation, ranging from occasional symptomatic nephrolithiasis to nephrocalcinosis and end-stage renal disease with systemic involvement. The prevalence is 11,000,000-	600,25
АНП	Joubert syndrome type 3	nm_001134830.1	NM_001134830.1:c.3263_3264delGG, NM_001134830.1:c.2295dupA, NM_001134830.1:c.21680>A, NM_001134830.1:c.1484G>A, NM_001134830.1:c.13030>T, NM_001134830.1:c.10520>T, NM_001134830.1:c.10510>T, NM_001134830.1:c.9850>T	9:1,000,000. Joubert syndrome type 3 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the AHII gene located on chromosomal region 6q25.3. The age of onset is variable. This disease is characterized by the neurological features of Joubert syndrome (neonatal hypotonia, developmental delay, mild to severe intellectrual disability, ataxia, and abnormal eye movements including oculomotor apraxia and primary position nystagmus) associated with retinal dystrophy.	600,25
AIPLI	Leber congenital amaurosis type 4	NM_014336.4	NM_014336.4:c.1053_1064delTGCAGAGCCACC, NM_014336.4:c.834G>A, NM_014336.4:c.715T>C, NM_014336.4:c.589G>C	amaurosis Type 4 (LCA4) is a severe dystrophy of the retina, typically becoming evident in the first years of life. Visual function is usually poor and often accompanied by nystagmus, sluggish or near-absent pupillary responses, photophobia, high hyperopia and keratoconus. Mutations in the AIPLI gene may cause approximately 20% of recessive LCA. Other conditions caused by pathogenic variants in the AIPLI gene mere conditions caused by pathogenic variants in the AIPLI gene mere conditions caused by pathogenic variants in the AIPLI gene are cone rod dystrophy and the less agressive form, juvenile retinitis pigmentosa. Cone-rod dystropy is characterized by decreased visual acuity, color vision defects, photoaversion and field, later followed by progressive loss in peripheral vision and night	600,25
ALDOB	Fructose intolerance, hereditary	NM_000035.3	NM_000035.3:c.1067C>A, NM_000035.3:c.1013C>T, NM_000035.3:c.1005C>C, NM_000035.3:c.720C>A, NM_000035.3:c.612T>A, NM_000035.3:c.612T>A, NM_000035.3:c.642T>C, NM_000035.3:c.442T>C, NM_000035.3:c.360_36364CAAA, NM_000035.3:c.178C>T, NM_000035.3:c.131_115delGGTA, NM_000035.3:c.10C>T, NM_000035.3:c.2T>C	blindness. Hereditary fructose intolerance follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ALDOB gene located on chromosomal region 9q21.3-q22.2. The age of onset is neonatal/infantile. This disease is characterized by severe abdominal pain, vomiting, and hypoglycemia following ingestion of fructose or other sugars metabolised through fructose-1-phosphate. The prevalence is 1:100,000-9:100,000. Congenital disorder of glycosylation type lc	
ALG6	Congenital disorder of glycosylation, type 1c	NM_013339.3	NM_013339.3:c.316C>T, NM_013339.3:c.897_899delAAT, NM_013339.3:c.998C>T, NM_013339.3:c.1432T>C	follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ALG6 gene located on chromosomal region 1p313. The age of onset is neonatal/infantile. This disease is characterized by psychomotor delay and muscular hypotonia, and possible coagulation anomalies, hormonal abnormalities and seizures. The prevalence is	600,25
ALMSI	Alstr ∲ m syndrome	NM_015120.4	NM_015120.4:c.2323C>T, NM_015120.4:c.4246delC, NM_015120.4:c.5584.6-T, NM_015120.4:c.8383C>T, NM_015120.4:c.9614_9618delCaCAA, NM_015120.4:c.11443C>T, NM_015120.4:c.11453dupA, NM_015120.4:c.11613.11613delCT, NM_015120.4:c.12439C>T, NM_015120.4:c.12445C>T	<1:1,000,000. Alstr	600,25

anos	Limb-girdle muscular dystrophy type 12 (LGMDR12; formerly LGMD2L)	NM_213599.2	NM_213599.2c:172C>T, NM_213599.2c:191dupA, NM_213599.2c:206_207delAT, NM_213599.2c:692G>T, NM_213599.2c:120C>T, NM_213599.2c:1295C>G, NM_213599.2c:1407+5G>A, NM_215599.2c:1627dupA, NM_213599.2c:1733T>C, NM_213599.2c:1887delA, NM_213599.2c:1898+1G>A, NM_213599.2c:1914G>A	is caused by pathogenic variants in the ANOS gene located on chromosomal region Tipl-43. This disease is characterized by weakness and wasting restricted to the limb musculature. Most often is characterized by an adult onset (but ranging from 11 to 51 years) of mainly proximal lower limb weakness, with difficulties standing on tiptoes being one of the initial signs. Proximal upper limb and distal lower limb weakness, is also common, as well as atrophy of the quadriceps (most commonly), biceps brachii, and lower leg muscles. Calf hypertrophy has also been reported in some cases. LCMDRI2 progresses slowly, with most patients remaining ambulatory until late adulthood. The estimated prevalence is <13,000,000.
АРТХ	Ataxia, early-onset, with oculomotor apraxia and hypoalbuminemia	NM_001195248.1	NM_001195248.1:c.917-1G>A, NM_001195248.1:c.879G>A, NM_001195248.1:c.830T>C, NM_001195248.1:c.659C>T, NM_001195248.1:c.1562delC, NM_001195248.1:c.166C>T NM_001195248.1:c.176-2A>G, NM_001195248.1:c.166C>T	Ataxia, early-onset, with oculomotor apraxia and hipoalbuminemia follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the APTX gene located on chromosomal region 9p31. Ataxia-oculomotor apraxia syndrome is an early-onset 600,25 autosomal recessive, progressive, cerebellar ataxia with peripheral axonal neuropathy, oculomotor apraxia (defined as the limitation of ocular movements on command), and hypoalbuminemia. The prevalence is unknown. The complete androgen insensitivity syndrome (CAIS) follows an X-linked pattern of inheritance and
AR	Androgen insensitivity syndrome, complete	NM_000044.3	NM_000044.3:c:340C>T, NM_000044.3:c:1771A>T, NM_000044.3:c:232SC>T, NM_000044.3:c:239IG>A, NM_000044.3:c:239SC>G, NM_000044.3:c:2567G>A, NM_000044.3:c:2650A>T	is caused by pathogenic variants in the AR gene located on chromosomal region Xq12. Affected males have female external genitalia, female breast development, blind vagina, absent uterus and female adnexa, and abdominal or inguinal testes, despite a normal male 46,7X karyotype. There is unresponsiveness to age-appropriate levels of androgens. There is also a partial androgen insensitivity syndrome (PAIS, OMIM 312300) caused by mutations in the AR gene, called Reifenstein syndrome, which results in hypospadias and micropenis with gynecomastia. Note: A specific type of mutation in the AR gene (a CAG repeat expansion) also cause a rare condition known as Spinal and
ARSA	Metachromatic leukodystrophy	NM_000487.5	NM_000487.5:c.1408_1418delGCAGCTGTGAC, NM_000487.5:c.1401_1411delGTTAGACGCAG, NM_000487.5:c.1283C>T, NM_000487.5:c.1241delC, NM_000487.5:c.1232C>T, NM_000487.5:c.1241delC, NM_000487.5:c.135C>A, NM_000487.5:c.134C>T, NM_000487.5:c.135C>A, NM_000487.5:c.134C>T, NM_000487.5:c.136C>A, NM_000487.5:c.936C>T, NM_000487.5:c.936C>A, NM_000487.5:c.937C>T, NM_000487.5:c.936C>A, NM_000487.5:c.893T>C, NM_000487.5:c.836C>A, NM_000487.5:c.893T>C, NM_000487.5:c.854C>A, NM_000487.5:c.893T>C, NM_000487.5:c.854C>A, NM_000487.5:c.827C>T, NM_000487.5:c.854C>A, NM_000487.5:c.827C>T, NM_000487.5:c.536C>A, NM_000487.5:c.545C>T, NM_000487.5:c.536C>A, NM_000487.5:c.545C>T, NM_000487.5:c.545C>T, CA, NM_000487.5:c.545C>T, NM_00487.5:c.465C>T, CA, NM_000487.5:c.545C>T, NM_00487.5:c.320C>A, NM_000487.5:c.346C>T, NM_00487.5:c.320C>A, NM_000487.5:c.325C>T, NM_000487.5:c.337G>A, NM_000487.5:c.325C>T, NM_000487.5:c.336C>A, NM_000487.5:c.395C>T, NM_000487.5:c.346elG	bulbar muscular atrophy or Kennedy disease; this mutation is not tested by this carrier test. Metachromatic leukodystrophy follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ARSA gene located on chromosomal region 22q13.33. The age of onset is variable. This disease is characterized by hypotonia, walking difficulties, optic atrophy and motor regression preceding mental impairment in the late infantile form, arrested intellectual development, followed by motor regression, epileptic seizures and ataxia in the juvenile form, and motor or psychiatric disorders, but with slow progression in the adult form. The incidence is 0.5:5,000-1:50,000 and the prevalence is 1:10,000 -5/10,000. Mucopolysaccharidosis type 6 (Maroteaux-Lamy) follows an autosomal recessive pattern of
			NM_000046.3:c.1438dupG, NM_000046.3:c.1366C>T, NM_000046.3:c.1124G>A, NM_000046.3:c.118A>C, NM_000046.3:c.1163-1G>C, NM_000046.3:c.1143-8T>G, NM_000046.3:c.979C>T,	inheritance and is caused by pathogenic variants in the ARSB gene located on chromosomal region 59(4.1. The age of onset is infantile. This lysosomal storage disorder resulting from a deficiency of arylsulfatase B is characterized by educed pulmonary function,

ARSB	mucopoiysaccnaridosis type 6 (Maroteaux-Lamy)	NM_0000463	NM_000046.3c.971G>T, NM_000046.3c.9244G>A, NM_000046.3c.937C>G, NM_000046.3c.937delA, NM_000046.3c.937C>G, NM_000046.3c.629A>G, NM_000046.3c.589C>T, NM_000046.3c.37C>T, NM_000046.3c.349T>C	nepatospienomegajy, hearing loss, sleep apnea, corneal clouding, carpal tunnel disease and occasionally central nervous system findings may include cervical cord compression caused by cervical spinal instability, meningeal thickening and/or bony stenosis, communicating hydrocephalus, optic nerve atrophy and blindness. The prevalence is 125,000-1600,000 newborns. X-linked chondrodysplasia punctata follows an X-linked pattern of inheritance and is caused by pathogenic variants in the APSE gene located on chromosomal region Xp2.33. The age of onset is neonatal. This is a disorder of cartilage and bone development that occurs almost exclusively in males. Include short stature and unusually short fingertips and ends of the toes. This condition is also associated with distinctive facial features, particularly a flattenedappearing nose with crescent-shaped nostrils and a flat nasal bridge.	600,25
ARSE	Chondrodysplasia punctata, X-linked recessive	NM_001282628.1	NM_001282628.1c.1807C>T, NM_001282628.1c.1517C>T, NM_001282628.1c.1504deIG, NM_001282628.1c.485C>T, NM_001282628.1c.194T>G, NM_001282628.1c.99-IG>A	People with X-linked chondrodysplasia punctata I typically have normal intelligence and a normal life expectancy. However, some affected individuals have had serious or life-threatening complications including abnormal thickening (stenosis) of the cartilage that makes up the airways, which restricts breathing. Also, abnormalities of spinal bones in the neck can lead to pinching (compression) of the spinal cord, which can cause pain, numbness, and weakness. Other, less common features of X-linked chondrodysplasia punctata I include delayed development, hearing loss, vision abnormalities, and heart defects. The prevalence is 1:500,000. Argininosuccinic aciduria follows an autosomal recessive pattern of inheritance and is caused	
ASL	Argininosuccinic aciduria	NM_000048.3	NM_000048.3c.35G>A, NM_000048.3c.337C>T, NM_000048.3c.346C>T, NM_000048.3c.346C>T, NM_000048.3c.446E1G>A, NM_000048.3c.532G>A, NM_000048.3c.532G>A, NM_000048.3c.532F3C, NM_000048.3c.532F3C, NM_000048.3c.532F3C, NM_000048.3c.532F3C, NM_000048.3c.532F3C, NM_000048.3c.6502F3C, NM_000048.3c.6502F3C, NM_000048.3c.1055C-T, NM_000048.3c.1056C-T, NM_000048.3c.1056C-T, NM_000048.3c.1056C-T, NM_000048.3c.1155C-T, NM_000048.3c.1255_1256delCT, NM_000048.3c.1369dupG	by pathogenic variants in the ASL gene located on chromosomal region 7q11.21. The age of onset is infantile. This disease is characterized by severe hyperammonemic coma, hypotonia, growth failure, anorexia and chronic vomiting or behavioral disorders during childhood, and hyperammonemic coma or behavioral disorders that simulate psychiatric disorders later in life. The prevalence is 170,000 newborns. Canavan disease follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ASPA gene located on chromosomal region 17p13.3. The age of	600,25
ASPA	Canavan disease	NM_000049.2	NM_000049.2:c.212G>A, NM_000049.2:c.433-2A>G, NM_000049.2:c.654C>A, NM_000049.2:c.633C>A, NM_000049.2:c.854A>C, NM_000049.2:c.914C>A	onset is neonatal/infantile. This disease is characterized by a variable spectrum between severe forms with leukodystrophy, macrocephaly and severe developmental delay, and a very rare mild/juvenile form characterized by mild developmental delay. The prevalence is 16,400- 113,500 in Askenazis Jewis.	
ASPM	Primary microcephaly type 5, autosomal recessive	³ NM_018136.4	NM_018136.4:c:10059C>A, NM_018136.4:c:9789T>A, NM_018136.4:c:9754delA, NM_018136.4:c:9742-9748delCT, NM_018136.4:c:9742-9748delCT, NM_018136.4:c:9697C>T, NM_018136.4:c:9697C>T, NM_018136.4:c:9697C>T, NM_018136.4:c:9697C>T, NM_018136.4:c:9697C>T, NM_018136.4:c:9697C>T, NM_018136.4:c:919C>T, NM_018136.4:c:919C=I, NM_018136.4:c:919	Primary autosomal recessive microcephaly type 5 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ASPM gene located on chromosomal region 1q31. The age of onset is neonatal/infantile. This disease is characterized by a reduction in head circumference at birth, mild to moderate non-progressive intellectual impairment and delay in early motor milestones, speech delay and	600,25

NM_DI8136.4:c.51881>G, NM_DI8136.4:c.5082G>A, NM_DI8136.4:c.3055C>T, NM_DI8136.4:c.2967G>A, NM_DI8136.4:c.2365C>T, NM_DI8136.4:c.2967G>A, NM_DI8136.4:c.2389C>T, NM_DI8136.4:c.1959.1962delCAAA, NM_DI8136.4:c.1729_1730delAG, NM_DI8136.4:c.1726_1730delAG, NM_DI8136.4:c.1350delAG, NM_DI8136.4:c.1350

hyperactive behavior are common. The annual incidence is 1:1,000,000.

			NM_018136.4:c.1002delA, NM_018136.4:c.719_720delCT, NM_018136.4:c.577C>T, NM_018136.4:c.349C>T		
			NM_000050.4:c.40G>A, NM_000050.4:c.256C>T, NM_000050.4:c.257C>A, NM_000050.4:c.349G>A, NM_000050.4:c.421- 2A>G, NM_000050.4:c.370G>A, NM_000050.4:c.396-2A>G, NM_000050.4:c.5375-C, NM_000050.4:c.396-SA, NM_000050.4:c.5371G>A, NM_000050.4:c.787G>A,	located on chromosomal region 9q34.1. The age of	
ASSI	Citrullinemia type 1	NM_000050.4	NM_000050.4'c.793C>T, NM_000050.4'c.794C>A, NM_000050.4'c.805C>A, NM_000050.4'c.814C>T, NM_000050.4'c.835C>T, NM_000050.4'c.836C>A, NM_000050.4'c.910C>T, NM_000050.4'c.939C>T, NM_000050.4'c.970C>A, NM_000050.4'c.970-5G>A, NM_000050.4'c.1085G>T, NM_000050.4'c.1087C>T, NM_000050.4'c.1088G>A, NM_000050.4'c.1086C>A,	onset is variable. This disease is characterized by hyperammonemia, progressive lethargy, poor feeding and vomiting in the neonatal form and by variable hyperammonemia in the	600,25
			NM_000050.4:c.1194-1G>C	later-onset form. The prevalence is 1:100,000- 9:100,000. AICA-ribosiduria due to ATIC deficiency follows an autosomal recessive pattern of inheritance and	
ATIC	AICA-ribosiduria due to ATIC deficiency	NM_004044.6	NM_004044.6:c.1277A>G	is caused by pathogenic variants in the ATIC gene located on chromosomal region 2q35. The age of onset is neonatal/infantile. This disease is characterized by profound intellectual deficit, epilepsy, dysmorphic features of the knees, elbows, and shoulders and congenital blindness. The prevalence is <13,00,000.	
АТР7В	Wilson disease	NM_000053.3	NM_000053.3c.4088C>T, NM_000053.3c.4058C>A, NM_000053.3c.3995C>T, NM_000053.3c.3995.2c, NM_000053.3c.3995C>T, NM_000053.3c.3995C>T, NM_000053.3c.3595C>T, NM_000053.3c.3595C>T, NM_000053.3c.3595T>A, NM_000053.3c.3595T>A, NM_000053.3c.3595T>A, NM_000053.3c.2597C>T, NM_000053.3c.2975C>T, NM_000053.3c.2975C>T, NM_000053.3c.2975C>T, NM_000053.3c.295C=A, NM_000053.3c.293C>T, NM_000053.3c.295C>T, NM_000053.3c.2795C>T, NM_000053.3c.2795C>T, NM_000053.3c.2795C>T, NM_000053.3c.2795C>T, NM_000053.3c.2795C>T, NM_000053.3c.2795C>T, NM_000053.3c.2795C>T, NM_000053.3c.2795C>T, NM_000053.3c.2795C>T, NM_00053.3c.2795C>T, NM_00053.3c.2795C>T, NM_000053.3c.2795C>T, NM_00053.3c.2795C>T, NM_00053.3c.2795C>T, NM_00053.3c.2795C>T, NM_00053.3c.2795C>T, NM_00053.3c.2795C>T, NM_000053.3c.2795C>T, NM_00053.3c.2795C>T,	Wilson disease follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ATP7B gene located on chromosomal region 13a(1-3. The age of onset is infantile. This disease is characterized by the toxic accumulation of copper, mainly in the liver and central nervous system, and symptomatic patients may present with hepatic, neurologic or psychiatric forms. The birth incidence is 130,000-1100,000 in	600,25
			NM_000053.3:c:19_20delCA	France and The prevalence is 1:10,000-1:30,000. Seckel syndrome type 1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ATR gene located on chromosomal region 3q23.	
ATR	Seckel syndrome type 1	NM_001184.3	NM_001184.3:c.6488delT, NM_001184.3:c.6037dupA, NM_001184.3:c.5645delA, NM_001184.3:c.5635Go-T, NM_001184.3:c.2341+1G>A, NM_001184.3:c.975_976delCT	The age of onset is neonatal/infantile. This disease is characterized by a proportionate dwarfism of prenatal onset, a severe microcephaly with a birdheaded like appearance and mental retardation. The prevalence is <1:1,000,000. Maple syrup urine disease type 1A follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the BCKDHA	600,25
BCKDHA	Maple syrup urine disease, type la	NM_000709.3	NM_000709.3:c.14delT, NM_000709.3:c.632C>T, NM_000709.3:c.659C>T, NM_000709.3:c.74ldupT, NM_000709.3:c.797delA, NM_000709.3:c.853C>C, NM_000709.3:c.9968C>A, NM_000709.3:c.954S>C, NM_000709.3:c.909.910delT, NM_000709.3:c.97delT, NM_000709.3:c.993C>C, NM_000709.3:c.964C>T, NM_000709.3:c.979C>A, NM_000709.3:c.1036C>T, NM_000709.3:c.1037C>A, NM_000709.3:c.1234G>A	gene located on chromosomal region 19q13.1-13.2. The age of onset is neonatal/infantile. This disease is characterized by poor feeding, lethargy, vomiting, a maple syrup odor in the cerumen and urine, encephalopathy and central respiratory failure if untreated. The prevalence is 11,000,000-11,000,000. Leigh syndrome caused by mutations in the BCSIL gene-located on chromosomal region 2q35-follows an autosomal recessive pattern of inheritance. Leigh syndrome is a clinically and genetically heterogeneous disorder resulting from defective mitochondrial energy generation; it presents extensive genetic heterogeneity (more than 75 different genes) with	600,25
BCSIL	BCS1L-related disorders, including Leigh syndrome	NM_001079866.1	NM_001079866.1c.103G>C, NM_001079866.1c.133C>T, NM_001079866.1c.148A>C, NM_001079866.1c.166C>T, NM_001079866.1c.232A>G, NM_001079866.1c.547C>T, NM_001079866.1c.548C>A, NM_001079866.1c.550C>T, NM_001079866.1c.5956d*T, NM_001079866.1c.590C>A, NM_001079866.1c.590C>A, NM_001079866.1c.590C>A	mutations identified in both nuclear- and mitochondrial-encoded genes involved in energy metabolism, including mitochondrial respiratory chain complexes I, II, III, IV, and V. It most commonly presents as a progressive and severe neurodegenerative disorder with onset within the first months or years of life, and may result in early	

				usually show global developmental delay or developmental regression, hypotonia, ataxia, dystonia, and ophthalmologic abnormalities, such as nystagmus or optic atrophy. The BCSIL protein is critical for the formation of mitochondrial complex III. This syndrome affects at least 1 in 40,000 newborns.	
BESTI	Bestrophinopathy, AR	NM_001139443.1	NM_001139443.1:c.242G>A, NM_001139443.1:c.341_342delTG, NM_001139443.1:c.348C>T, NM_001139443.1:c.348C>T, NM_001139443.1:c.502C>A, NM_001139443.1:c.769G>A, NM_00139443.1:c.769G>A, NM_00139443.1:c.1129_1130insCCAAAGA, NM_00139443.1:c.1203_1204insGCCTTGATGGA, NM_001139443.1:c.1311_1317dupCAAAGAC	Bestrophinopathy follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the BESTI gene located on chromosomal region liql3. The age of onset is variable. This disease is characterized by central visual loss in the first 2 decades of life associated with an absent electrocculogram light rise, and a reduced electroretinogram. Genetic heterogeneity: Mutations in this gene may cause dominant phenotypes like Macular dystrophy, vitelliform, 2 (OMIM 153700) and Vitreoretinochoroidopathy (195220).	600,25
BESTI	Bestrophinopathy, AR	NM_004183.3	NM_004183.3:c.122T>C	an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the BESTI gene located on chromosomal region 1(q15. The age of onset is variable. This disease is characterized by central visual loss in the first 2 decades of life associated with an absent electrooculogram light rise, and a reduced electroretinogram. Genetic heterogeneity. Mutations in this gene may cause dominant phenotypes like Macular dystrophy, vitelliform, 2 (600,25
BSND	Bartter syndrome, type 4a	NM_057176.2	NM_057176.2:c.1A>T, NM_057176.2:c.3G>A, NM_057176.2:c.10G>T, NM_057176.2:c.22G>T, NM_057176.2:c.35T>C, NM_057176.2:c.139G>A	OMIM 153700) and Vitreoretinochroriodpathy (193220). Bartter syndrome type 4A with deafness follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the BSND gene located on chromosomal region 1p32.3. The age of onset is neonatal/infantile. This disease is characterized by maternal polyhydramnios, premature delivery, polyuria, sensorineural deafness and is associated with hypokalemic alkalosis, increased levels of plasma renin and	600,25
BTD	Biotinidase deficiency	NM_001281723.2	NM_001281723.2c.1900>A, NM_001281723.2c.241C>T, NM_001281723.2c.340C>C, NM_001281723.2c.440C>T, NM_001281723.2c.544C>T, NM_001281723.2c.534G>T, NM_001281723.2c.534G>T, NM_001281723.2c.534G>T, NM_001281723.2c.5354C, NM_001281723.2c.635A<0, NM_001281723.2c.635C>T, NM_001281723.2c.635A<0, NM_001281723.2c.637C>T, NM_001281723.2c.649C>T, NM_001281723.2c.649C>T, NM_001281723.2c.6396deIT, NM_001281723.2c.1374S-C, NM_001281723.2c.1374S-C, NM_001281723.2c.1356C>T, NM_001281723.2c.1356C>T	aldosterone, low blood pressure, and vascular resistance to angiotensin II. Biotinidase deficiency an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the BTD gene located on chromosomal region 3p25. The age of onset is neonatal/infantile. This disease is characterized by seizures, breathing difficulties, hypotonia, skin rash, alopecia, hearing loss and delayed development. Limb-girdle muscular dystrophy type 1 (LCMDR1; formerly LCMDR2), formerly LCMDR2, formerly LCMDR2, formerly LCMDR2, formerly LCMDR2, formerly LCMDR2, formerly LCMDR3, former	600,25
CAPN3	Limb-girdle muscular dystrophy type 1 (LGMDR1; formerly LGMD2A)	NM_000070.2	NM_000070.2:c.133G>A, NM_000070.2:c.223dupT, NM_000070.2:c.328C>T, NM_000070.2:c.550delA, NM_000070.2:c.530elF, NM_000070.2:c.550delT, NM_000070.2:c.550delT, NM_000070.2:c.550delT, NM_000070.2:c.550delT, NM_000070.2:c.1550delT, NM_000070.2:c.1550delT, NM_000070.2:c.1566C>T, NM_000070.2:c.1569C>T, NM_000070.2:c.1569C>T, NM_000070.2:c.1559_1602delGAGC, NM_000070.2:c.1575G>A, NM_000070.2:c.1559_1602delGAGC, NM_000070.2:c.1550delAGGC, NM_000070.2:c.1550delAGGC, NM_000070.2:c.1550delAGGC, NM_000070.2:c.1550delAGGC, NM_000070.2:c.1550delAGGC, NM_000070.2:c.1550delAGGC, NM_000070.2:c.1550delAGGC, NM_000070.2:c.236delAGGCC, NM_000070.2:c.236delAGGCC, NM_000070.2:c.236delAGGCC, NM_000070.2:c.236delAGGCCC, NM_000070.2:c.236delAGGCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	an autosomal recessive pattern of inheritance and is caused by biallelic pathogenic variants in the CAPNS gene located on chromosomal region 15q15.1. The age of onset is variable. This disease is characterized by a variable age of onset of progressive, typically symmetrical and selective weakness and atrophy of proximal shoulder- and pelvic-girdle muscles (gluteus maximus, thigh adductors, and muscles of the posterior compartment of the limbs are most commonly affected) without cardiac or facial involvement. Clinical manifestations include exercise intolerance, a waddling agit, scapular winging and calf pseudo-hypertrophy.The prevalence is 1:10,000-9:100,000. Genetic heterogeneity.	

death. Affected individuals usually show global

				the CAPN3 gene can cause autosomal dominant limb-girdle muscular dystrophy-4 (LGMDD4; OMIM 618129),	
CBS	Homocystinuria, B6- responsive and nonresponsive types	NM_000071.2	NM_000071.2c.1330G>A, NM_000071.2c.1280C>T, NM_000071.2c.1150A>G, NM_000071.2c.1036G>A, NM_000071.2c.1036G>A, NM_000071.2c.1036G>T, NM_000071.2c.1036G>A, NM_000071.2c.1036G>A, NM_000071.2c.995G>A, NM_000071.2c.995G>A, NM_000071.2c.995G>A, NM_000071.2c.935T>C, NM_000071.2c.975G>A, NM_000071.2c.575G>A, NM_000071.2c.575G>A, NM_000071.2c.576G>T, NM_000071.2c.576G>T, NM_000071.2c.576G>T, NM_000071.2c.576G>T, NM_000071.2c.576G>T, NM_000071.2c.576G>T, NM_000071.2c.576G>T, NM_000071.2c.576G>T, NM_000071.2c.576G>T, NM_000071.2c.376G>A, NM_000071.2c.376G>T, N	prevalence is 1.200,000- 1.335,000. Joubert syndrome type 9 defect follows an autosomal recessive pattern of inheritance and	600,25
CC2D2A	Joubert syndrome type 9; Meckel syndrome type 6	NM_001080522.2	NM_001080522.2:c.2486+1G>C, NM_001080522.2:c.2848C>T, NM_001080522.2:c.345C>T, NM_001080522.2:c.345C>T, NM_001080522.2:c.3369delG, NM_001080522.2:c.3369-41G>A, NM_001080522.2:c.345C>T, NM_001080522.2:c.4191-41delG, NM_001080522.2:c.4191-41delG, NM_001080522.2:c.4582C>T, NM_001080522.2:c.4582C>T, NM_001080522.2:c.4582C>T, NM_001080522.2:c.4667A>T	is caused by pathogenic variants in the CC2D2A gene located on chromosomal region 4pl5.32. The age of onset is neonatal/infantile. This disease is characterized neonatal hypotonia, developmental delay, intellectrual disability, ataxia, and abnormal eye movements including oculomotor apraxia, primary position, primary position systagmus and congenital	600,25
CDH23	Usher syndrome, type 1D	NM_022124.5	NM_022124.5:c.146-2A>G, NM_022124.5:c.193delC, NM_022124.5:c.1858+2T>G, NM_022124.5:c.1858+2T>G, NM_022124.5:c.3162.5*14C>A, NM_022124.5:c.5316_3.5*1964ATCC, NM_022124.5:c.556_33T>G, NM_022124.5:c.5634C>T, NM_022124.5:c.5633T>G, NM_022124.5:c.5633T>G, NM_022124.5:c.5633T>G, NM_022124.5:c.6050-9G>A, NM_022124.5:c.6393delC, NM_022124.5:c.6442G>A	10p22.1. The age of onset is neonatal/infantile. This disease is characterized by congenital, non- progressive, mild-to- profound sensorineural	600,25
CDHRI	Cone-rod dystrophy, type 15	NM_033100.3	NM_033100.3:c.338delG, NM_033100.3:c.524dupA, NM_033100.3:c.640delG, NM_033100.3:c.1112delC, NM_033100.3:c.1485+2T>C, NM_033100.3:c.1485+2T>G	hearing impairment. Cone-rod dystrophy, type 15 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CDHRI gene located on chromosomal region 10q231. This disease is characterized by decreased visual acuity and sensitivity in the central visual field, followed by loss of peripheral vision. The overall prevalence of all	600,25
CENPJ	Primary microcephaly type 6, autosomal recessive	° NM_018451.4	NM_018451.4:c.3842_3843dupTA, NM_018451.4:c.3704A>T, NM_018451.4:c.3699_3702dupAATA, NM_018451.4:c.3568_3.571dupCTCA, NM_018451.4:c.3415G>T, NM_018451.4:c.3243_3246deITCAG, NM_018451.4:c.26162.2972deIAAAAA, NM_018451.4:c.2614deIT, NM_018451.4:c.260_2463deICACG, NM_018451.4:c.1959_1952dupAGTG NM_018451.4:c.757_760deIGTCT, NM_018451.4:c.1949_1952dupAGTG NM_018451.4:c.322_236deICAGAA, NM_018451.4:c.40C>T	types of cone-rod dystrophy is 1-9:100,000. Primary autosomal recessive microcephaly type 6 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CENPJ gene located on chromosomal region 13q12.12. The age of onset, is neonatal/infantile. This disease is characterized by reduced head circumference at birth with no gross anomalies of brain architecture and	600,25
CEP290	Meckel syndrome type 4; Joubert syndrome type 5	NM_025114.3	NM_025114.3c;7341dupA, NM_025114.3c;7341delA, NM_025114.3c;7341delA, NM_025114.3c;7324G>T, NM_025114.3c;6798G>A, NM_025114.3c;6645+1G>A, NM_025114.3c;6624delG, NM_025114.3c;6645+1G>A, NM_025114.3c;6516564delCAA, NM_025114.3c;4962,4963delAA, NM_025114.3c;4962,4963delAA, NM_025114.3c;4962,4963delAA, NM_025114.3c;4962,4963delAA, NM_025114.3c;4962,4963delAA, NM_025114.3c;4962,4963delA, NM_025114.3c;4930S>T, NM_025114.3c;185delT, NM_025114.3c;1652,1665delAA, NM_025114.3c;15016>T, NM_025114.3c;1616>T, NM_025114.3c	variable degrees of intellectual impairment. Meckel syndrome type 4 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CEP290 gene located on chromosomal region 12q21.32. The age of onset is neonatal. This disease is characterized by a combination of renal cysts and variably associated features including developmental anomalies of the central nervous system (typically occipital encephalocele), hepatic ductal dysplasia and cysts,	
CERKL	Retinitis pigmentosa type 26	NM_001030311.2	NM_001030311,2:c:1090C>T, NM_001030311.2:c:858delT, NM_001030311,2:c:847C>T, NM_001030311.2:c:312delA	and postaxial polydactyly. The prevalence is <1/1,000,000. Retinitis pigmentosa 26 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CERKL gene located on chromosomal region 29313. The age of onset is variable. This disease is characterized by night blindness (nyctalopia), peripheral visual field impairment and over time loss of central vision. The prevalence is 130,000-510,000. Complement factor H deficiency follows an	600,25

CEH

NM_000186.3:c.380G>T, NM_000186.3:c.1606T>C, NM_000186.3:c.2876G>A

autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CFH gene located on chromosomal region 1q32. This disease is characterized by increased 600.25 characterized by increased susceptibility to recurrent, usually severe, infections (particularly by Neisseria meningitidis, Escherichia coli, and Haemophilus influenza), repai influenzae), renal impairment and/or autoimmune diseases

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NM_000492.3:c:1A>G, NM_000492.3:c:4C>T, NM_000492.3:c:11C>A, NM_000492.3:c:504eIT, NM_000492.3:c:44T>C, NM_000492.3:c:53+1C>T, NM_000492.3:c:57C>A, NM_000492.3:c:57C>A, NM_000492.3:c:36C>T, NM_000492.3:c:15C>T, NM_000492.3:c:15C
                NM_000492.3c:79G-1, NM_000492.3c:88C-1, NM_0004

NM_000492.3c:137C-A, NM_000492.3c:164-16-A,

NM_000492.3c:164-16-T, NM_000492.3c:164-27-C,

NM_000492.3c:165-16-A, NM_000492.3c:165-3C-T,

NM_000492.3c:165-16-A, NM_000492.3c:170G-A,

NM_000492.3c:171G-A, NM_000492.3c:170G-A,

NM_000492.3c:171G-A, NM_000492.3c:178G-A,

NM_000492.3c:171G-A, NM_000492.3c:178G-A,

NM_000492.3c:173G-A, NM_000492.3c:178G-A,
NM_000492.3c:171G>A, NM_000492.3c:178_177delTAGA, NM_000492.3c:178G>T, NM_000492.3c:178G>T, NM_000492.3c:178G>T, NM_000492.3c:178G>T, NM_000492.3c:200C>T, NM_000492.3c:178G>T, NM_000492.3c:235dupT, NM_000492.3c:245dupT, NM_000492.3c:2535dupT, NM_000492.3c:2535dupT, NM_000492.3c:257dupT, NM_000492.3c:257dupT, NM_000492.3c:257dupT, NM_000492.3c:257dupT, NM_000492.3c:257dupT, NM_000492.3c:258dupT, NM_000492.3c:258dupT
          NM_000492.3c:.803delA, NM_000492.3c:.825C>G, NM_000492.3c:.828C>A, NM_000492.3c:.828C>A, NM_000492.3c:.828C>A, NM_000492.3c:.83CabdupA, NM_000492.3c:.835_937delTCT, NM_000492.3c:.935_937delTCT, NM_000492.3c:.935_937delTCT, NM_000492.3c:.936CabdupA, NM_000492.3c:.936CabdupA, NM_000492.3c:.908CabdupA, NM_000492.3c:.010G>T, NM_0004
          NM_000492.3:c.1355_1156dupTA, NM_000492.3:c.1202G>A, NM_000492.3:c.1203G>A, NM_000492.3:c.1203G>A, NM_000492.3:c.1209+1G>A, NM_000492.3:c.1240G>T, NM_000492.3:c.1240G>T, NM_000492.3:c.1240G>T, NM_000492.3:c.13501_1307delCaCTTCT, NM_000492.3:c.13501_1307delCaCTTCT, NM_000492.3:c.1350dupCaTA, NM_000492.3:c.1340delA, NM_000492.3:c.1353CaC, NM_000492.3:c.1397G>A, NM_000492.3:c.1397G>A, NM_000492.3:c.1397G>A, NM_000492.3:c.1397G>A, NM_000492.3:c.1436G=C, NM_000492.3:c.1436G=C, NM_000492.3:c.1475G>T, NM_000492.3:c.1475G>T, NM_000492.3:c.1477G>T, NM_000492.3:c.1477G>T, NM_000492.3:c.1475G>T, NM_000492.3:c.1475G>T, NM_000492.3:c.1475G>T, NM_000492.3:c.1457G>A, NM_000492.3:c.1457G>A, NM_000492.3:c.1551=156A>G, NM_000492.3:c.15151=1525delCTT, NM_000492.3:c.151524=6G-G, NM_000492.3:c.151521=525delCTT,
NM_000492.3c.;1519_1521de1ATC, NM_000492.3c.;1556T>C, NM_000492.3c.;1519_1521de1ATC, NM_000492.3c.;1516A>G, NM_000492.3c.;1519_1527de1ATC, NM_000492.3c.;1516A>G, NM_000492.3c.;1519_1523de1CTT, NM_000492.3c.;1519_1523de1CTT, NM_000492.3c.;1519_1523de1CTT, NM_000492.3c.;1519_1523de1CTT, NM_000492.3c.;1519_1523de1ATC, NM_000492.3c.;1586A>G, NM_000492.3c.;1587A>G, NM_000492.3c.;1786A>G, NM_000492.3c.;1786A>AC, NM_000492.3c.;1786AAAACTA, NM_000492.3c.;1786AAACTA, NM_000492.3c.;1786AAAACTA, NM_000492.3c.;1786AAACTA, NM_000492.3c.;1
                NM_000492.3c.2053dupC, NM_000492.3c.2053C>T,
NM_000492.3c.21265A*T, NM_000492.3c.2128A*T,
NM_000492.3c.21243C>T, NM_000492.3c.2138C>T,
NM_000492.3c.2135dupA, NM_000492.3c.2139ST>G,
NM_000492.3c.2215deIG, NM_000492.3c.2242248deIGATACTGC,
NM_000492.3c.2290C>T, NM_000492.3c.2353C>T,
NM_000492.3c.2290C>T, NM_000492.3c.2435C>T,
NM_000492.3c.2374C>T, NM_000492.3c.2435C>T,
NM_000492.3c.2453deIT, NM_000492.3c.2436.2464deITG,
NM_000492.3c.2453deIT, NM_000492.3c.2463.2464deITG,
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Cystic fibrosis follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CFTR gene located on chromosoma region 7q31.2. The age of onset of severe form is neonatal or infantile but there are also variants associated with moderate clinical or late onset. This disease is characterized by the production of sweat with a high salt content, mucus secretions with an abnormal viscosity, chronic bronchitis pancreatic insufficiency adolescent diabetes and more rarely, stercoral obstruction and cirrhosis. Male sterility is a constant feature. Late-onset forms, which are usually only mild or monosymptomatic. The prevalence is 1:10,000 9:10,000.

CETP

Cystic fibrosis

NM_000492.3

NM_000492.3:c.2988+1G>A, NM_000492.3:c.2989-2A>G, NM_000492.3:c.2989-1G>A, NM_000492.3:c.3002_3003delTG,

NM_000492.3c.3013.019delCTATAGCAG, NM_000492.3c.3017C>A, NM_000492.3c.3039dupC, NM_000492.3c.3039delC, NM_000492.3c.3039delC, NM_000492.3c.3039delC, NM_000492.3c.30367_3072delATAGTG, NM_000492.3c.3080T>C, NM_000492.3c.31392.3c.31810=26.2c.31810=2 NM_000492.3c.32095-A, NM_000492.3c.32050-A, NM_000492.3c.32050-A, NM_000492.3c.3276C-A, NM_000492.3c.3276C-A, NM_000492.3c.3276C-A, NM_000492.3c.3276C-A, NM_000492.3c.3276C-A, NM_000492.3c.32924C-A, NM_000492.3c.32924C-A, NM_000492.3c.32924C-A, NM_000492.3c.32924C-T, NM_000492.3c.32924C-T, NM_000492.3c.32924C-T, NM_000492.3c.32924C-T, NM_000492.3c.3302T-A, NM_000492.3c.3300C-T, NM_000492.3c.3350C-T, NM_000492.3c.3550C-T, NM_000492.3c.35 Macular corneal dystrophy follows an autosomal recessive pattern of recessive pattern of inheritance and is caused by pathogenic variants in the CHST6 gene located on chromosomal region on the ChST6. Space of the CHST6 are provided on chromosomal region on the CHST6 are provided by the CHST6. Macular corneal dystrophy NM_021615.4 NM 021615.4:c.327 328delC1 variable. This disease is variable. Inis disease is characterized by bilateral ill-defined cloudy regions within a hazy stroma, and eventually severe visual impairment. The prevalence is 1:100,000-9:100,000. Myotonia congenita follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CLCN1 gene located on chromosomal region 7q35. The age of onset is neonatal/infantile. This is a nondystrophic skeletal muscle disorder NM 000083.2:c.180+3A>T, NM 000083.2:c.225dupC. muscle disorder characterized by muscle stiffness and an inability of 600,25 the muscle to relax after voluntary contraction. Most patients have NM_000083.2:c.409T>G, NM_000083.2:c.871G>A, NM_000083.2:c.1238T>G, NM_000083.2:c.1453A>G, NM_000083.2:c.2680C>T Mvotonia congenita. CLCN1 NM_000083.2 symptom onset in the legs, which later progresses to the arms, neck, and facial muscles. Many patients show marked hypertrophy of the lower limb muscles The prevalence is 1:100.000. Renal hypomagnesemia type 5, with ocular involvement follows an autosomal recessive pattern of inheritance and is caused by pathogeni variants in the CLDN19 gene located on Rena hypomagnesemia NM_148960.2:c.425_437delCCCTGGTGACCCA, NM_148960.2:c.269T>C, NM_148960.2:c.169C>G, NM_148960.2:c.59G>A chromosomal region CLDN19 NM_148960.2 type 5, with ocular 1p34.2. The age of onset is 600,25 infantile. This disease is volvement infantile. This disease is characterized by excessive magnesium and calcium renal wasting, bilateral nephrocalcinosis, progressive renal failure and severe ocular abnormalities. The prevalence is <1:1,000,000. Usher syndrome type 3A follows an autosoma follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CLRN1 gene located on chromosomal region. NM_001195794.1:c.669_670insT, NM_001195794.1:c.630dupT, 3q25.1. The age of onset is neonatal/infantile. This CLRN1 Usher syndrome, type 3A NM 001195794.1 NM_001195794.1:c.189C>A, NM_001195794.1:c.144T>G, NM_001195794.1:c.18T>G, NM_001195794.1:c.92C>T 600.25 disease is characterized by the association of sensorineural deafness with retinitis pigmentosa and progressive vision loss. The prevalence is 1:1.000.000- 9/1.000.000. Retinitis pigmentosa 49 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CNGAI gene located on chromosomal region 4p12. The age of onset is variable. This disease is NM_001142564.1:c.2179delA, NM_001142564.1:c.2134C>T, NM_001142564.1:c.11747C>T, NM_001142564.1:c.1166C>T, NM_001142564.1:c.1001G>A, NM_001142564.1:c.656-2T>C, NM_001142564.1:c.304dupA Retinitis pigmentosa type NM_001142564.1

characterized by night

CNGB1	Retinitis pigmentosa type 45	NM_001297.4	NM_001297.4:c.3462+1G>A, NM_001297.4:c.3425deIT. NM_001297.4:c.3150deIG, NM_001297.4:c.2762.2765deIACGA, NM_001297.4:c.12653deIG, NM_001297.4:c.1249.2475-G, NM_001297.4:c.1958-1G>A, NM_001297.4:c.1122-2A>T, NM_001297.4:c.952C-T, NM_001297.4:d.315-1G>A, NM_001297.4:c.218-	characterized by night	600,25
			2A>G NM_019098.4:c.2048_2049delCA, NM_019098.4:c.2011G>T,	blindness, perjpheral visual field impairment and over time loss of central vision. The prevalence is 1:10,000 to 5:10,000. Achromatopsia type 3 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CNGB3 gene located on chromosomal region 8q213. The age of onset is neonatal/Infantile. This disease is characterized by reduced visual acuity,	
CNGB3	Achromatopsia type 3	NM_019098.4	NM_019098.4:c.1148delC, NM_019098.4:c.1063C>T, NM_019098.4:c.893_897delCAAAA, NM_019098.4:c.897_896delCTTCTACAAA, NM_019098.4:c.896_890delACTTC, NM_019098.4:c.819_826delCAGACTCC, NM_019098.4:c.446_447insT	pendular nystagmus, increased sensitivity to light (photophobia), a small central scotoma, and reduced or complete loss of color discrimination. Most individuals have complete form, with total lack of function in all three types of cones. Rarely, individuals have incomplete form, with similar, but generally less severe symptoms. The prevalence is 1/30,000-1/50,000. Epidermolysis bullosa, junctional, non-Herlitz	
COL17A1	Epidermolysis bullosa, junctional, non-Herlitz type	NM_000494,3	NM_000494.3:c.4319dupC, NM_000494.3:c.4003_4004delGG, NM_000494.3:c.3998G>A, NM_000494.3:c.3897_3900delATCT, NM_000494.3:c.3897_3900delATCT, NM_000494.3:c.3876-1, NM_000494.3:c.3975delC, NM_000494.3:c.30576-7, NM_000494.3:c.3042-7, NM_000494.3:c.30576-7, NM_000494.3:c.3042-7, NM_000494.3:c.2954-1, NM_000494.3:c.2954-1, NM_000494.3:c.2954-1, NM_000494.3:c.2554-7, NM_000494.3:c.2536-7, NM_000494	type follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the COLT/Al gene located on chromosomal region 10q24-3. The age of onset is neonatal/infantile. This disease is characterized by a generalized skin bilstering, atrophic scarring, nail dystrophy or nail absence, and enamel hypoplasia, with extracutaneous involvement. Knobloch syndrome type 1 follows an autosomal	600,25
COLIBAI	Knobloch syndrome, type 1	NM_130444.2	NM_130444_2:c.1700_1701insGACGTGAAAGAGGGG, NM_130444_2:c.2240_224iinsGACGTGAAAGAGGGG, NM_130444_2:c.3994_3295delAQ, NM_130444_2:c.3502C>T, NM_130444_2:c.4072_4084delCCCCCAGGCCCAC, NM_130444_2:c.4214_4223delCAGGGCCCCC, NM_130444_2:c.4222_4225delCC, NM_130444_2:c.4323_4323+1delGG, NM_130444_2:c.4272_4759_4760delCT, NM_130444_2:c.5168dupG	recessive pattern of inheritance and is caused by pathogenic variants in the COLIBAI gene located on chromosomal region 2lq22.3. The age of onset is neonatal/infantile. This disease is characterized by vitreoretinal and macular degeneration, and occipital encephalocele. The prevalence is <11.000.000. Alport syndrome, autosomal recessive follows an autosomal recessive inheritance and is caused inheritance and is caused	600,25
COL4A3	Alport syndrome, autosomal recessive	NM_000091.4	NM_000091.4:c.345delG, NM_000091.4:c.898G>A, NM_000091.4:c.2083G>A, NM_000091.4:c.211delC, NM_000091.4:c.2954G>T, NM_000091.4:c.4420_442delCTTTT, NM_000091.4:c.4441C>T, NM_000091.4:c.457IC>G	by pathogenic variants in the COL4A4 and COL4A4 genes located on chromosomal region 2,365.3 The age of onset is infantile. This disease is characterized by renal, cochlear, and ocular involvement. Renal disease progresses from microscopic hematuria to proteinuria, progressive renal insufficiency, and end-stage renal disease. Progressive sensorineural hearing loss is usually present by late childhood or early adolescence. Ocular findings include anterior lemticomus, maculopathy, corneal endothelial vesicles, and recurrent corneal erosion. The prevalence is 1:50,000 newborn. Alport syndrome, autosomal recessive follows an autosomal recessive follows an autosomal recessive inheritance and is caused	600,25
COL4A4	Alport syndrome, autosomal recessive	NM_000092.4	NM_000092.4:c.4923C>A, NM_000092.4:c.4129C>T, NM_000092.4:c.3713C>A, NM_000092.4:c.3601G>A, NM_000092.4:c.71+1G>A	by pathogenic variants in the COL4A3 and COL4A4 genes located on chromosomal region 2363. The age of onset is infantile. This disease is characterized by renal, cochlear, and ocular involvement. Renal disease progresses from	600,25

				Progressive sensormedral hearing loss is usually present by late childhood or early adolescence. Ocular findings include anterior lenticonus, maculopathy, corneal endothelial vesicles, and recurrent corneal erosion. The prevalence is 1:50,000 newborn.	
COL7A1	Epidermolysis bullosa dystrophica, AR	NM_000094.3	NM_000094.3:c.8524_8527+10delGAAGGTCAGGACAG, NM_000094.3:c.84795-T, NM_000094.3:c.84206-T, NM_000094.3:c.8393T>A, NM_000094.3:c.8245G>A, NM_000094.3:c.7957G>A, NM_000094.3:c.8245G>A, NM_000094.3:c.7957G>A, NM_000094.3:c.79350-TG>C, NM_000094.3:c.7952G-T, NM_000094.3:c.6946G>A, NM_000094.3:c.685G-A, NM_000094.3:c.6573-TG>T, NM_000094.3:c.6857G>A, NM_000094.3:c.6573-TG>T, NM_000094.3:c.6570G>T, NM_000094.3:c.6573-TG>T, NM_000094.3:c.652TdupC, NM_000094.3:c.6573-TG>T, NM_000094.3:c.582T-TG>T, NM_000094.3:c.5532+TG>A, NM_000094.3:c.582T-TG>T, NM_000094.3:c.5532+TG>A, NM_000094.3:c.582T-TG>T, NM_000094.3:c.503G>T, NM_000094.3:c.582T-TG>T, NM_000094.3:c.4888C>T, NM_000094.3:c.582T-TG>T, NM_000094.3:c.4783C>T, NM_000094.3:c.4783G>C, NM_00094.3:c.4704G>T, NM_000094.3:c.4783G>C, NM_00094.3:c.471dupG, NM_000094.3:c.4781G>T, NM_00094.3:c.8471dupG, NM_000094.3:c.953G>C, NM_000094.3:c.8471dupG, NM_000094.3:c.953G>C, NM_000094.3:c.8471dupG, NM_000094.3:c.953G>C, NM_000094.3:c.8471dupG, NM_000094.3:c.756C>T, NM_000094.3:c.8576elG, NM_000094.3:c.756C>T, NM_000094.3:c.8576elG, NM_000094.3:c.356C>G	Epidermolysis bullosa dystrophica follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the COL7A1 gene located on chromosomal region 3p21. The age of onset is neonatal/infantile. This disease is characterized by generalized cutaneous and mucosal bilistering and scarring associated with severe deformities and major extracutaneous involvement. The prevalence is <13,000,000.	600,25
COQ2	Primary coenzyme Q10 deficiency, type 1	NM_015697.7	NM_015697.7:c.1197delT, NM_015697.7:c.890A>G, NM_015697.7:c.723delT NM_015697.7:c.683A>G, NM_015697.7:c.590G>A	Primary coenzyme Q10 deficiency type 1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the COQ2 gene located on chromosomal region 4q21.23. The age of onset is neonatal/infantile. The phenotypes include an encephalomyopathic form with seizures and ataxia; a multisystem infantile form with encephalopathy, cardiomyopathy and renal failure; a predominantly cerebellar form with ataxia and cerebellar atrophy; Leigh syndrome with growth retardation; and an isolated myopathic form.	600,25
COQ8A	Primary coenzyme Q10 deficiency, type 4	NM_020247.4	NM_020247.4:c.589-3C>G, NM_020247.4:c.637C>T, NM_020247.4:c.815G>A, NM_020247.4:c.815G>T, NM_020247.4:c.911C>T, NM_020247.4:c.1541a>G, NM_020247.4:c.1759_1752delACC, NM_020247.4:c.1813dupG	Primary coenzyme Q10 deficiency type 4 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the COQ8A gene located on chromosomal region lq42.13. The age of onset is infantile. This disease is characterized by progressive ataxia, cerebellar atrophy, and often exercise intolerance with elevated lactate levels and mild intellectual deficit.	600,25
СРТ2	Carnitine palmitoyltransferase type 2 deficiency, lethal neonatal	NM_000098.2	NM_000098.2:c.149C>A, NM_000098.2:c.338C>T, NM_000098.2:c.359A>G, NM_000098.2:c.370C>T, NM_000098.2:c.520G>A, NM_000098.2:c.520G>A, NM_000098.2:c.520G>A, NM_000098.2:c.520G>A, NM_000098.2:c.525.725edelAC, NM_000098.2:c.520S, NM_000098.2:c.137C>T, NM_000098.2:c.137C>T, NM_000098.2:c.137C>T, NM_000098.2:c.137C>T, NM_000098.2:c.137C>T, NM_000098.2:c.137C>T, NM_000098.2:c.1383A>C, NM_000098.2:c.1381C>T	Carnitine palmitoyltransferase deficiency, type 2, lethal neonatal form follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CPT2 gene located on chromosomal region 1p32. The age of onset is neonatal/infantile. This disease is characterized by a severe fasting intolerance leading to metabolic derangements of hypoketotic hypoglycemia, resulting in coma and seizures, and hepatic encephalopathy leading to liver failure. The prevalence is <1:,000,000.	
CRB1	Retinitis pigmentosa type 12, AR; Leber congenital amaurosis type 8	NM_201253.2	NM_201253.2:c.498_506delAATTGATGG, NM_201253.2:c.613_619delATAGGAA, NM_201253.2:c.2290C>T, NM_201253.2:c.2618T_A, NM_201253.2:c.246G>T, NM_201253.2:c.2688T_A, NM_201253.2:c.2983C>T, NM_201253.2:c.3055_3059dupTATAT, NM_201253.2:c.3122T>C, NM_201253.2:c.3383delT, NM_201253.2:c.3299T>C, NM_201253.2:c.3389delT, NM_201253.2:c.3419T>A, NM_201253.2:c.3999TG>T	Retinitis pigmentosa type I2 and leber congenital amaurosis type 8 follow an autosomal recessive pattern of inheritance and are caused by pathogenic variants in the CRBI gene located on chromosomal region lgal-ag2.1. Retinitis pigmentosa type 12 is characterized by night blindness, peripheral visual field impairment and over time loss of central visionm, and its prevalence is 1-5:10,000. Leber congenital amaurosis, with a neonatal/infantile age of onset, comprises a group of early-onset childhood retinal dystrophies characterized by vision loss, nystagmus, and severe retinal dysfunction. Patients usually present at birth with profound vision loss and pendular nystagmus. Other clinical findings of this disease may include high hypermetropia, photodysphoria, oculodigital sign, keratoconus, cataracts, and a variable appearance to the fundus. Nephropathic cystinosis follows an autosomal recessive pattern of	600,25

CTNS	Cystinosis (atypical/juvenile/ocular) nephropathic	NM_001031681.2	NM_0010316812:c.283G>T, NM_0010316812:c.329G>T, NM_0010316812:c.357_360delCAGC, NM_0010316812:c.397_398delAT, NM_0010316812:c.36C>T, NM_0010316812:c.36C>T, NM_0010316812:c.59G>A, NM_0010316812:c.59G>A, NM_0010316812:c.5853-3C>G, NM_0010316812:c.1015G>A	inheritance and is caused by pathogenic variants in the CTNS gene located on chromosomal region 17pl3. The age of onset is neonatal/infantile. This disease is characterized by hypothyroidism, insulindependent diabetes, hepatosplenomegaly with portal hypertension, and muscle, cerebral and ocular involvement, caused by cystine deposits in various organs. The prevalence is 1:100,000-1200,000.	
стѕк	Pycnodysostosis	NM_000396.3	NM_000396.3:c.926T>C, NM_000396.3:c.721C>T, NM_000396.3:c.436G>C, NM_000396.3:c.236G>A, NM_000396.3:c.154A>T	Pycnodysostosis follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CTSk gene located on chromosomal region Iq21. The age of onset is variable. This disease is characterized by osteosclerosis, short stature or dwarfism, acroosteolysis of the distal phalanges, fragile bones associated with spontaneous fractures and dysplasia of the clavicles. The prevalence is 11,000,000 to 91,000,000.	600,25
CYP4V2	Bietti crystalline corneoretinal dystrophy	NM_207352.3	NM_207352.3:c.130T>A, NM_207352.3:c.327+1G>A, NM_207352.3:c.332T>C, NM_207352.3:c.1523G>A	Bietti crystalline corneoretinal dystrophy follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CYP4V2 gene located on chromosomal region 4q35.2. The age of onset is adult. This disease is characterized by nightbilmdness, decreased vision, paracentral scotoma, and, in the end stages of the disease, legal blindness. Spastic paraplegia type 5A	600,25
СҮР7В1	Spastic paraplegia type SA autosomal recessive	NM_004820.4	NM_004820.4:c.1460dupT, NM_004820.4:c.1456C>T, NM_004820.4:c.18562>T, NM_004820.4:c.889A>G, NM_004820.4:c.187C>T	follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CYP7B1 gene located on chromosomal region 8q213. The age of onset is neonatal/infantile. This disease is characterized by a slow, gradual, progressive weakness and spasticity of the lower limbs. Rate of progression and the severity of	600,25
D2HGDH	D-2-hydroxyglutaric aciduria	NM_152783.4	NM_152783.4:c.440T>G, NM_152783.4:c.1123G>T, NM_152783.4:c.1315A>G, NM_152783.4:c.1331T>C, NM_152783.4:c.1333_1334delAC	D-2-Hydroxyglutaric aciduria follows autosomal recessive pattern of inheritance and is caused by pathogenic variants in the D2HGDH gene located on chromosomal region 2q37.3. The age of onset is variable. This disease is characterized by extremely variable clinical, manifestations, with severe cases characterized by neonatal or early infantile-onset epileptic encephalopathy, and marked hypotonia, and cerebral visual failure, developmental delay, seizures, involuntary movements, and cardiomyopathy are also common in these cases.	600,25
DBT	Maple syrup urine disease, type 2	NM_001918.3	NM_001918.3:c.1281+1G>A, NM_001918.3:c.939G>C, NM_001918.3:c.901C>T, NM_001918.3:c.871C>T, NM_001918.3:c.827T>G, NM_001918.3:c.772+1G>A, NM_001918.3:c.570C>T, NM_001918.3:c.581C>G, NM_001918.3:c.294C>G, NM_001918.3:c.272_275delCAGT, NM_001918.3:c.126T>G	The prevalence is below 1,000,000. Maple syrup urine disease, type 2 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the DBT gene located on chromosomal region 1p21.2. The age of onset in neonatal/infantil. This disease is characterized by a maple syrup odor to the urine, deficient diet, lethargy and focal dystonia, followed by progressive encephalopathy and central respiratory failure if not treated. The prevalence is 1-5/10,000. Omenn syndrome and Athabascan type severe combined immunodeficiency follow	600,25

pattern of inheritance and are caused by pathogenic variants in the DCLREIC gene located on chromosomal region 10p13. Omenn syndrome has an early age of onset and it is characterized by desquamation, alopecia, chronic diarrhea, failure to thrive, lymphadenopathy, Omenn syndrome; Severe combined immunodeficiency, NM_001033855.2:c.1639G>T, NM_001033855.2:c.1558dupA, NM_001033855.2:c.597C>A, NM_001033855.2:c.597C>A, NM_001033855.2:c.2T>C Athabascan type and hepatosplenomegaly associated with severe combined combined immunodeficiency. The age of onset of Athabascan type severe combined immunodeficiency is neonatal/infantile and it is a combined in the second immunodeficiency is neonatal/infantile and it is neonatal/infantile and it characterized by severe and recurrent infections, diarrhea, failure to thrive, and cell sensitivity to ionizing radiation. The prevalence is 1-9/1,000,000. Mitochondrial DNA depletion syndrome type 3 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the DGUOK gene located on chromosomal region 2p13. The age of onset is neonatal/infantile. This NM_080916.2:c.137A>G, NM_080916.2:c.313C>T, NM_080916.2:c.425G>A, NM_080916.2:c.494A>T, NM_080916.2:c.707+2T>G, NM_080916.2:c.763G>T DGUOK-related mitochondrial DNA NM_080916.2 depletion syndrome disease is characterized by disease is characterized by progressive liver failure, hypoglycemia and neurologic abnormalities including hypotonia, encephalopathy and propagative and propagative control of the contro peripheral neuropathy Smith-Lemli-Opitz syndrome follows an autosomal recessive NM_001163817.1:c.1342G>A, NM_001163817.1:c.1337G>A, NM_001163817.1:c.1228G>A, NM_001163817.1:c.1210C>T, NM_001163817.1:c.1055G>A, NM_001163817.1:c.1054C>T, pattern of inheritance and pattern of inheritance and is caused by pathogenic variants in the DHCR7 gene located on chromosomal region 11q13.4. The age of onset is 600,25 neonatal/infantile. This disease is characterized by NM_0011638177:e.1055G>A, NM_0011638177:e.1054G>T, NM_0011638177:e.907G>T, NM_0011638177:e.907G>T, NM_0011638177:e.907G>T, NM_0011638177:e.907G>A, NM_0011638177:e.907G>A, NM_0011638177:e.707G>T, NM_0011638177:e.707G>T, NM_0011638177:e.707G>T, NM_0011638177:e.725G>A, NM_0011638177:e.725G>A, NM_0011638177:e.725G>A, NM_0011638177:e.725G>A, NM_0011638177:e.725G>A, NM_0011638177:e.725G>A, NM_0011638177:e.725G>A, NM_0011638177:e.735G>T, NM_0011638177:e.735G Smith-Lemli-Opitz NM 001163817.1 syndrome disease is characterized by onsease is characterized by multiple congenital anomalies, intellectual deficit, and behavioral problems. The prevalence is 1/20,000 to 1/40,000 newborn. Dihydropyrimidine dehydrogenase deficiency follows an autosomal recessive pattern of recessive pattern of inheritance and is caused by pathogenic variants in the DPYD gene located on chromosomal region 1p22. This disease shows large phenotypic variability, pnenotypic variability, ranging from no symptoms to a convulsive disorder with motor and mental retardation in homozygous patients. In people with severe dihydropyrimidine dehydrogenase deficiency, the disorder becomes the disorder becomes apparent in infancy. These affected individuals have recurrent seizures (epilepsy), intellectual disability, a small head size Dihydropyrimidine dehydrogenase deficiency NM_000110.3 NM_000110.3:c.1905+1G>A, NM_000110.3:c.1679T>G, NM_000110.3:c.1109_1110delTA, NM_000110.3:c.299_302delTCAT DPVD disability, a small head size (microcephaly), increased muscle tone (hypertonia), delayed development of motor skills such as walking, and autistic behaviors that affect communication and social interaction. The interaction. The prevalence is unknow. In addition, homozygous and heterozygous mutation carriers can develop severe toxicity after the administration of the antineoplastic drug 5fluorouracil (5FU). Dilated cardiomyopathy with woolly hair and keratoderma, known as Keratoderma, known as Carvajal syndrome, follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the DSP gene located on chromosomal located on chromosomal region 6p24. The age of onset is neonatal/infantile. This disease is characterized by woolly hair is present at birth and the palmoplantar keratoderma appears keratoderma appears during the first year of life. The cardiac anomaly presents during childhood and is marked by dilation of the left ventricle accompanied by accompanied by alterations in muscle contractility. The dilated cardiomyopathy may lead to life-threatening congestive heart failure and death. The prevalence is below 1,000,000. Cardiomyopathy, dilated NM 004415 3:c 3098del4 NM 004415 3:c 5800C>T DSP NM 004415.3

with woolly hair and keratoderma; Epidermolysis bullosa lethal acantholytic

NM_004415.3:c.6370_6371delCT, NM_004415.3:c.7000C>T NM_004415.3:c.7180_7181delAG, NM_004415.3:c.8188C>T

Furthermore, mutations in

pattern of inheritance and

the DSP gene have been identified in people with an autosomal recessive disorder called lethal acantholytic epidermolysis bullosa. Features of this bullosa. Features of this condition include very fragile skin that blisters and detaches easily, a complete absence of hair (alopecia), abnormal or missing fingernalis, teeth that are present from birth that are present from birth (neonatal teeth), and abnormalities of the heart muscle (cardiomyopathy). The skin abnormalities lead to a severe loss of fluids and death in early infancy. Miyoshi muscular dystrophy, type 1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the DYSF gene located on chromosomal region 2p13.3. The age of

onset is young adulthood. This disease is This disease is characterized by weakness and atrophy in the distal lower extremity posterior compartment (gastrocnemius and soleus muscles) and is associated with difficulties.

associated with difficulties associated with difficulties in standing on tip toes.
The prevalence is 1/1,000,000 to 9/1,000,000. 600,25 Mutations in the DYSF gene can also cause muscular dystrophy, limb-

girdle, autosomal

girdle, autosomal recessive, type 2. This disease is characterized by an onset in late adolescence or early adulthood of slowly progressive, proximal weakness and atrophy of shoulder and pelvic girdle shoulder and pelvic girdle muscles. Cardiac and respiratory muscles are not involved. Hypertrophy of the calf muscles and highly elevated serum creatine kinase levels are

frequently observed. Miyoshi muscular dystrophy, type 1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the DYSF gene located on chromosomal region 2p13.3. The age of onset is young adulthood This disease is

This disease is characterized by weakness and atrophy in the distal lower extremity posterior compartment (gastrocnemius and soleus muscles) and is associated with difficulties

associated with difficulties in standing on tip toes. The prevalence is 1/1,000,000 to 9/1,000,000. Mutations in the DYSF gene can also cause muscular dystrophy, limbgirdle, autosomal

girdle, autosomal recessive, type 2. This disease is characterized by an onset in late adolescence or early adulthood of slowly progressive, proximal weakness and atrophy of weakness and atrophy of shoulder and pelvic girdle muscles. Cardiac and respiratory muscles are not involved. Hypertrophy of the calf muscles and highly elevated serum creatine kinase levels are

frequently observed. Hypohidrotic ectodermal hyponiarotic ectodermal dysplasia, type 1, hypohidrotic, X-linked policy of inheritance and is caused by pathogenic variants in the EDA gene located on chromosomal

600.25

region Xq12-q13.1. The age of onset is neonatal/infantile. This disease is characterized by malformation of ectodermal structures such as skin, hair, teeth and sweat glands. The prevalence is 1/5,000 to

1/10,000 newborns Trichothiodystrophy (TTD), type 1 is a heterogeneous type I is a heterogeneous group of disorders that follows an autosomal recessive pattern of inheritance. It is caused by pathogenic variants in the ERCC2 gene located on chromosomal region

chromosomal region 19q13.32. The age of onset is neonatal or infantile. This disease, with variable clinical expression, is characterized by brittle and fragile hair, often combined with growth

Mivoshi muscular dystrophy, type 1; Muscular NM_001130978.1 NM_001130978.1:c.1481-1G>A autosomal recessive, type 2 DYSE

> NM_001130987.1c.203_204delTGinsAT,
> NM_001130987.1c.203_204delTGinsAT,
> NM_001130987.1c.396_397delCC, NM_001130987.1c.706C>T,
> NM_001130987.1c.359-1G>C, NM_001130987.1c.797G>A,
> NM_001130987.1c.391G>T, NM_001130987.1c.1033-1G>A,
> NM_001130987.1c.1149+1G>A, NM_001130987.1c.1033-1G>A,
> NM_001130987.1c.1360-27-C, NM_001130987.1c.16464C>A,
> NM_001130987.1c.1360-27-C, NM_001130987.1c.1649-2A>C,
> NM_001130987.1c.1694-1C-A, NM_001130987.1c.1692-27-A,
> NM_001130987.1c.1674delA, NM_001130987.1c.1692-27-A,
> NM_001130987.1c.177C>T, NM_001130987.1c.1692-27-A,
> NM_001130987.1c.177C>T, NM_001130987.1c.1827C>T,
> NM_001130987.1c.1886C>T, NM_001130987.1c.1927C>T,
> NM_001130987.1c.2924_2928delAGACC, NM_00130987.1c.2923C>T,
> NM_001130987.1c.2924_274.1C.10130987.1c.30987.1c.2923C>T,
> NM_001130987.1c.2016.7 NM_00130987.1c.30987.1c.2923C>T,
> NM_001130987.1c.2016.7 NM_00130987.1c.30987.1c.2923C>T,

Ectodermal dysplasia, type NM_001399.4 EDA 1, hypohidrotic, X-linked

Miyoshi muscular

DYSF

dystrophy, type 1; Muscular NM_001130987.1 dystrophy, limb-girdle, autosomal recessive, type 2

NM_001399.4:c.181T>C, NM_001399.4:c.183C>G, NM_001399.4:c.187G>A, NM_001399.4:c.463C>T, NM_001399.4:c.466C>T, NM_001399.4:c.467C>A, NM_001399.4:c.573_574insT, NM_001399.4:c.575_574insT, NM_001599.4:c.575_575_574insT, NM_001599.4:c.575_575_575_575_575_575_575_575_5

NM_000400.3;c,2230_2233dupCTAG, NM_000400.3;c,2176C>T. NM_000400.3c: 22450_2253dupc | IAG, NM_000400.3c: 2176 NM_0004003c: 20470-7; NM_000400.3c: 19720-7; NM_000400.3c: 1703_1704delTT, NM_000400.3c: 1621A>C, NM_000400.3c: 135462-7; NM_000400.3c: 1381C>G, NM_000400.3c: 135462-7; NM_000400.3c: 13081-G>A, NM_000400.3c: 195-10-3c, NM_000400.3c: 5967G>A, NM_000400.3c: 719-10-3c, NM_000400.3c: 567G>A, NM_000400.3c: 179-10-3c, NM_000400.3c: 567G>A,

Trichothiodystrophy, type 1 NM_000400.3

ERCC5	Cerebrooculofacioskeletal syndrome, type 3	NM_000123.3	NM_000123.3:c.88+2T>C, NM_000123.3:c.215C>A, NM_000123.3:c.381-2A>G, NM_000123.3:c.406C>T, NM_000123.3:c.4664dupA, NM_000123.3:c.526C>T, NM_000123.3:c.275C>T, NM_000123.3:c.275T>CT, NM_000123.3:c.2575T>C, NM_000123.3:c.255T>C, NM_000123.3:c.25T>C, NM_000123.3:c.255T>C, NM_000	retaruation and intellectual deficit, congenital ichthyosis and nail abnormalities, among other symptoms. About half of the patients with TTD exhibit marked photosensitivity. Cerebrooculofacioskeletal syndrome type 3, also known as COFS syndrome, follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ERCCS gene located on chromosomal region 13g33.1. COFS syndrome is characterized by prenatal onset of arthrogryposis, microcephaly and growth failure. Postnatal features include severe developmental delay, congenital cataracts (in some), and marked UV sensitivity of the skin. Survival beyond 6 years of age is rare. The prevalence is below 1/1,000,000. Cockayne syndrome (CS), type B follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ERCC6 gene located on	600,25
ERCC6	Cockayne syndrome, type B; Cerebrooculofacioskeletal syndrome, type 1	NM_000124.3	NM_000124.3:c.3862C>T, NM_000124.3:c.3591_3592dupGA, NM_000124.3:c.2503C>T, NM_000124.3:c.2503C>T, NM_000124.3:c.2503C>T, NM_000124.3:c.2047C>T, NM_000124.3:c.1550C>A, NM_000124.3:c.357C>T, NM_000124.3:c.422+1C>A, NM_000124.3:c.207dupG, NM_000124.3:c.48_49delCT	chromosomal region 10q11.23. The age of onset is variable. This disease is characterized by short stature, a characteristic facial appearance, premature aging, photosensitivity, progressive neurological dysfunction, and intellectual deficit. Mutations in the ERRCG gene have been also found in patients with COFS syndrome type 1, an extreme prenatal form of the CS clinical spectrum. This autosomal recessive progressive neurodegenerative disorder is characterized	600,25
EYS	Retinitis pigmentosa, type 25	nm_001292009.1	NM_0012920091:c.9468T>A, NM_0012920091:c.9362_9365delCTCA, NM_0012920091:c.9099delT, NM_0012920091:c.9012.8692_8695dupACAG, NM_0012920091:c.8632G>T, NM_0012920091:c.8692_8695dupACAG, NM_0012920091:c.8632G>T, NM_0012920091:c.78922C>T, NM_0012920091:c.78922C>T, NM_0012920091:c.78922C>T, NM_0012920091:c.78922C>T, NM_0012920091:c.78922C>T, NM_0012920091:c.5928-2A>G, NM_0012920091:c.5928-2A>G, NM_0012920091:c.58926-2A>G, NM_0012920091:c.5928-2A>G, NM_0012920091:c.4120C>T, NM_0012920091:c.4120C>T, NM_0012920091:c.571dupA, NM_0012920091:c.932delT, NM_0012920091:c.103C>T, NM_0012920091:c.322delT, NM_0012920091:c.103C>T	by microcephaly, congenital cataracts, severe mental retardation, facial dysmorphism, and arthrogryposis. Retinitis pigmentosa, type 25 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the EYS gene located on chromosomal region 6q12. The age of onset is variable. This disease is characterized by night blindness (nyctalopia), peripheral visual field impairment and over time loss of central vision. The prevalence is 1/10,000 to 5/10,000.	600,25
FII	Factor XI deficiency, autosomal recessive	NM_000128.3	NM_000128.3:c.166T>C, NM_000128.3:c.403G>T, NM_000128.3:c.438C>A, NM_000128.3:c.595+3A>G, NM_000128.3:c.901T>C, NM_000128.3:c.1211C>A, NM_000128.3:c.1613C>T, NM_000128.3:c.1693G>A	characterized by reduced levels and activity of factor XI resulting in moderate bleeding symptoms, usually occurring after trauma or surgery. The prevalence is 1/1,000,000	600,25
F9	Hemophilia B	NM_000133.3	NM_000133.3:c.82T>C, NM_000133.3:c.1031T>C, NM_000133.3:c.1136G>A, NM_000133.3:c.1150C>T	to 9/1,000,000. Hemophilia B follows an X-linked pattern of inheritance and is caused by pathogenic variants in the F9 gene located on chromosomal region Xq27.1-q27.2. The age of onset is neonatal/infantile. This disease is characterized by spontaneous or prolonged hemorrhages due to factor IX deficiency. The prevalence is 1/100,000 to 9/100,000. Tyrosinemia, type 1 follows	600,25
FAH	Tyrosinemia, type 1	NM_000137.2	NM_000137.2:c.47A>T, NM_000137.2:c.192G>T, NM_000137.2:c.401C>A, NM_000137.2:c.456G>A, NM_000137.2:c.554-1G>T, NM_000137.2:c.707-1G>A, NM_000137.2:c.78GG>A, NM_000137.2:c.78GG>A, NM_000137.2:c.8339delC, NM_000137.2:c.9320e1C, NM_000137.2:c.1090G>A, NM_000137.2:c.1027G>T, NM_00137.2:c.1090G>A, NM_00137.2:c.1069G>T, NM_00137.2:c.1069G>T, NM_00137.2:c.1069G>T, NM_00137.2:c.1069G>T, NM_00137.2:c.1069G>T, NM_00137.2:c.1041A>G	an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the FAH gene located on chromosomal region 15q251. The age of onset is variable. This disease is characterized by progressive liver disease.	

				Fanconi anemia, complementation group A follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in	
FANCA	Fanconi anemia, complementation group A	NM_000135.2	NM_000135.2c.4130C>G, NM_000135.2c.3788_3790deITCT, NM_000135.2c.3763G>T, NM_000135.2c.3558dupG, NM_000135.2c.2303T>C, NM_000135.2c.1115_1118deITTGG, NM_000135.2c.233_236deITTGA, NM_000135.2c.131dupA	characterized by progressive pancytopenia with bone marrow failure,	600,25
			NM_000136.2:c:1642C>T, NM_000136.2:c:1487T>G,	variable congenital malformations and predisposition to develop hematological or solid tumors. The prevalence is 1/1,000,000 to 9/1,000,000. Fanconi anemia, complementation group C follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the FANCC gene located on chromosomal region 9422.3. The age of onset is	
FANCC	Fanconi anemia, complementation group C	NM_000136.2	NM_000136.2:c.1103_1104delTG, NM_000136.2:c.1015delA, NM_000136.2:c.996+1G>T, NM_000136.2:c.67delG, NM_000136.2:c.37C>T	infantile. This disease is	600,25
FANCG	Fanconi anemia, complementation group G	NM_004629.1	NM_004629.1:c.1852_1853delAA, NM_004629.1:c.1795_1804delToGATCCGTC, NM_004629.1:c.1480+1G>C, NM_004629.1:c.1077-2A>C, NM_004629.1:c.907_908dupCT, NM_004629.1:c.637_643delTACCGCC, NM_004629.1:c.510+1G>A, NM_004629.1:c.313G>T	infantile. This disease is characterized by progressive pancytopenia with bone marrow failure, variable congenital malformations and predisposition to develop hematological or solid	600,25
FGB	Congenital afibrinogenemia	NM_005141.4	NM_005141.4:c.1148T>G, NM_005141.4:c.1289G>A	tumors. The prevalence is 1,0,00,000 to 9/1,000,000. Congenital afibrinogenemia follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the FGB gene located on chromosomal region 4q2B. The age of onset is variable. This disease is characterized by bleeding symptoms ranging from mild to severe resulting from reduced quantity and/or quality of circulating fibrinogen. The prevalence	600,25
				is 1/1,000,000 to 9/1,000,000. To 9/1,000,000. Charcot-Marie-Tooth disease, type 4J follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the FIG4 gene located on chromosomal region 6q.21. The age of onset is neonatal/infantile. This disease is characterized by rapidly progressive, asymmetric motor neuron degeneration with slow nerve conduction	
FIG4	Charcot-Marie-Tooth disease, type 43; Yunis- Varon syndrome	NM_014845.5	NM_014845.5:c:122T>C, NM_014845.5:c:31IG>A, NM_014845.5:c:50IC>G, NM_014845.5:c:592C>T, NM_014845.5:c:737G>A, NM_014845.5:c:592C>T, NM_014845.5:c:737G>A, NM_014845.5:c:831_838delTAAATTTG, NM_014845.5:c:2299dupG	4/100,000 to 8/100,000. Mutations in the FIG4 gene have been also found in patient with Yunis-Varon syndrome. This disease is a severe autosomal recessive disorder characterized by skeletal defects, including cleidocranial dysplasia and severe neurologic involvement with neuronal loss. Enlarged cytoplasmic vacuoles are found in neurons, muscle, and cartilage. The disorder is usually lethal in infancy. Muscular dystrophydystroglycanopathy type 5 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the FKRP gene located on chromosomal region 19q13.32. The age of onset is neonatal or early infancy. There are three subtypes of dystroglycanopathy there are three subtype SA, SB and SC.	
	Muscular dystrophy-		NIM MINZOGOE 2- 1600-T NIM MINZOGOE 2- 11640-A	severe phenotype and is associated with congenital brain and eye anomalies, cobblestone lissencephaly, professed mostal	

			NINE AND A POLICE AND DATE OF THE AND A POLICE AND A PARTY.	protouriu mentar	
FKRP	dystroglycanopathy, type 5A, 5B and 5C	NM_001039885.2	NM_001033063.2:c.1343C>T, NM_001033685.2:c.1387A>G	retardation, and death usually in the first years of life. Included diseases are the more severe Walker- Warburg syndrome and	600,25
				the slightly less severe muscle-eye-brain disease.	
				Subtype 5B represents an intermediate phenotype with or without congenital	
				mental retardation, white matter changes and	
				structural brain abnormalities. Finally,	
				subtype 5C is the less severe phenotype	
				characterized by limb- girdle muscular dystrophy, variable age at onset,	
				normal cognition, and no structural brain changes.	
				Fragile X syndrome follows an X-linked pattern of	
				inheritance and is caused by pathogenic variants in	
				the FMR1 gene located on chromosomal region Xq27.3. The symptoms are	
				variable depending on the range of CGG triplet	
				expansion. In complete mutation the onset is	
		2		infantile in men and is characterized by intellectual disability,	
FMR1	Fragile X syndrome	-0	(CGG)n pre-mutated allele	characteristic appearance (large head, long face,	600,25
				prominent forehead and chin, protruding ears) joint	
				laxity and large testes after puberty. In carrier female, the symptoms are milder	
				and include primary ovarian insufficiency. The	
				prevalence is 1/2,500 (full mutation allele) to 1/4,000	
				(prevalence of symptomatic cases) for both genders.	
				Fraser syndrome, type 1 follows an autosomal	
				recessive pattern of inheritance and is caused	
				by pathogenic variants in the gene FRASI located on chromosomal region	
			NM_025074.6:c.835_838delGTGT, NM_025074.6:c.3799C>T,	4q21.21. The age of onset is early infancy. Twenty-five	
FRAS1	Fraser syndrome, type 1	NM_025074.6	NM_025074.6:c.5605_5606insT, NM_025074.6:c.6433C>T, NM_025074.6:c.6991_6992insGG, NM_025074.6:c.7813C>T, NM_025074.6:c.11160_11167deIGCTGGAGA	per cent of affected infants are stillborn, while	600,25
				20 % die before the age of 1 year. This disease is characterized mainly by	
				cryptophthalmos and syndactyly, besides urinary	
				and genital anormalities. The prevalence is	
				<1:1,000,000. Glycogen storage disease,	
				type 1A follows an autosomal recessive pattern of inheritance and	
			NM_000151.3:c.113A>T, NM_000151.3:c.229T>C, NM_000151.3:c.230+1G>C, NM_000151.3:c.247C>T, NM_000151.3:c.248G>A, NM_000151.3:c.370G>A,	is caused by pathogenic variants in the G6PC gene	
G6PC	Glycogen storage disease, type 1A	NM_000151.3	NM_000151.3:c.247621; NM_000151.3:c.24862A; NM_000151.3:c.379_380dupTA, NM_000151.3:c.447-1G>A, NM_000151.3:c.497T>C, NM_000151.3:c.508C>T, NM_000151.3:c.562G>C,	region 17q21.31. The age of	600,25
			NM_000151.3:c.883C>T, NM_000151.3:c.1039C>T	onset is infantile. This disease is characterized by poor tolerance to fasting,	
				significant hepatomegaly and growth retardation.	
				The incidence is 1/100,000. Glycogen storage disease,	
				type 1A follows an autosomal recessive pattern of inheritance and	
				is caused by pathogenic variants in the G6PC gene	
G6PC	Glycogen storage disease, type 1A	NM_001270397.1	NM_001270397.1:c,474G>A	located on chromosomal region 17q21.31. The age of	600,25
				onset is infantile. This disease is characterized by poor tolerance to fasting,	
				significant hepatomegaly and growth retardation.	
				The incidence is 1/100,000. Krabbe disease follows an	
				autosomal recessive pattern of inheritance and is caused by pathogenic	
			NM_000153.3:c.2056T>C, NM_000153.3:c.1964delC, NM_000153.3:c.1814dupA, NM_000153.3:c.1796T>G,	variants in the GALC gene located on chromosomal	
			NM_000153.3:c.1723.1724:insT, NM_000153.3:c.1700A>C, NM_000153.3:c.1695delT, NM_000153.3:c.1592G>A, NM_000153.3:c.1591C>T, NM_000153.3:c.1586C>T,	region 14q31.3. There are two forms of the disease:	
GALC	Krabbe disease	NM_000153.3	NM_000153.3:c.1543G>A, NM_000153.3:c.1489+1_1489+2delGT, NM_000153.3:c.1488_1489+2delTGGT, NM_000153.3:c.1488_1489delTG,	infantile form (2-6 months onset) more severe and adult form less severe. It is	600,25
			NM_000153.3:c.1472delA, NM_000153.3:c.1161+2T>G, NM_000153.3:c.153G>T, NM_000153.3:c.658C>T, NM_000153.3:c.758C	a degenerative disorder that affects the nervous	
			NM_000153.3:c.582+1G>A, NM_000153.3:c.453G>A, NM_000153.3:c.453G>A, NM_000153.3:c.453G>A,	system characterized by a muscle stiffness,	
			NM_000153.3:c.205C>T	blindness, deafness, and eventually death. The incidence is 1/100,000-	
				1/250,000 and the prevalence is 1/100,000.	
			NM_000155.3:c.18delC, NM_000155.3:c.41delCinsTT, NM_000155.3:c.71_72insA, NM_000155.3:c.135A>C,		
			NM_000155.3:c.118G>T, NM_000155.3:c.130G>A, NM_000155.3:c.132delG, NM_000155.3:c.152G>A, NM_000155.3:c.158G>A, NM_000155.3:c.199C>T, NM_000155.3:c.203A>C, NM_000155.3:c.220_221delCT,		
			NM_000155.3:c.221T>C, NM_000155.3:c.253-2A>G, NM_000155.3:c.265T>G, NM_000155.3:c.289_291delAAC,	Galactosemia follows an	
			NM_000155.3:c.290A>G, NM_000155.3:c.292G>A, NM_000155.3:c.329- 2A>C, NM_000155.3:c.367C>T, NM_000155.3:c.386T>C,	autosomal recessive pattern of inheritance and	
			NM_000155.3:c.400delT, NM_000155.3:c.404C>T, NM_000155.3:c.413C>T, NM_000155.3:c.425T>A, NM_000155.3:c.428C>T, NM_000155.3:c.428C>T, NM_000155.3:c.426C>T, NM_000155.3:c.42C>T, NM_000155.3:c.42C	variants in the GALT gene located on chromosomal	
			NM_000155.3:c.443G>A, NM_000155.3:c.502_504delGTG, NM_000155.3:c.505C>A, NM_000155.3:c.508-1G>C,	region 9p13.3. The age of onset is neonatal. This	
			NM_000155.3:c.512T>C, NM_000155.3:c.547C>A, NM_000155.3:c.552C>A,	alsease is characterized by	

GALT	Galactosemia	NM_000155.3	NM_000155.3°c.563A>G, NM_000155.3°c.565_578delGTATGGGCCAGCAG, NM_000155.3°c.568T>C, NM_000155.3°c.596T>C, NM_000155.3°c.584T>C, NM_000155.3°c.598delC, NM_000155.3°c.601C>T, NM_000155.3°c.602C>A, NM_000155.3°c.607G>A, NM_000155.3°c.603C>A, NM_000155.3°c.689C>A>C, NM_000155.3°c.634C>T, NM_000155.3°c.688-2A>C, NM_000155.3°c.634C>T, NM_000155.3°c.688-2A>C, NM_000155.3°c.772C>T, NM_000155.3°c.775C>T, NM_000155.3°c.790delC, NM_000155.3°c.772C>T, NM_000155.3°c.795C>T, NM_000155.3°c.790delC, NM_000155.3°c.774C>T, NM_000155.3°c.795C>T, NM_000155.3°c.790delC, NM_000155.3°c.794C>T, NM_000155.3°c.875C>T, NM_000155.3°c.790delC, NM_000155.3°c.844C>C, NM_000155.3°c.855C>T,	disease. Long-term complications appear including cognitive impairments, motor deficits, and ovarian dysfunction with reduced	
			NM_000155.3c:904-1C>T, NM_000155.3c:905-2A>G, NM_000155.3c:939C>A, NM_000155.3c:947C>A, NM_000155.3c:957C>A, NM_000155.3c:985T>C, NM_000155.3c:998C>A, NM_000155.3c:97C>T, NM_000155.3c:998C>A, NM_000155.3c:1006A>T, NM_000155.3c:1030C>A, NM_000155.3c:1048delA, NM_000155.3c:1052delC, NM_000155.3c:1138T>C	Giant axonal neuropathy, type I follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the GAN gene located on chromosomal region 16q23.2. The age of	
GAN	Giant axonal neuropathy, type 1	NM_022041.3	NM_022041.3:c.12681>C, NM_022041.3:c.1429C>T, NM_022041.3:c.1447C>T, NM_022041.3:c.1456G>A	disease is characterized by a progressive motor and sensitive peripheral and central nervous system neuropathy. Twenty families have been reported with this disease but the frequency is likely to be under-estimated. Gaucher disease follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the GBA gene located on chromosomal region 1q22. Gaucher disease encompasses a continuum of clinical findings from a perinatal lethal disorder to an asymptomatic type. There are three major clinical types (1, 2, and 3) and two other subtypes (perinatal-lethal and cardiovascular). Type 1 is characterized by the presence of clinical or radiographic evidence of bone disease,	600,25
GBA	Gaucher disease	NM_000157.3	NM_000157.3c.1604G-A, NM_000157.3c.1504C-T, NM_000157.3c.1448T-C, NM_000157.3c.1448T-C, NM_000157.3c.1448T-C, NM_000157.3c.1448T-C, NM_000157.3c.1348T-A, NM_000157.3c.1348T-A, NM_000157.3c.1346T-C, NM_000157.3c.1309G-T, NM_000157.3c.1309G-T, NM_000157.3c.1309G-T, NM_000157.3c.1309G-T, NM_000157.3c.1295G-T, NM_000157.3c.1295G-T, NM_000157.3c.1246G-A, NM_000157.3c.124G-S-A, NM_000157.3c.124G-S-A, NM_000157.3c.124G-S-C, NM_000157.3c.134G-S-C, NM_000157.3c.134G-S-C, NM_000157.3c.134G-S-C, NM_000157.3c.134G-S-C, NM_000157.3c.134G-S-C, NM_000157.3c.134G-S-C, NM_000157.3c.135G-S-T, NM_000157.3c.134G-S-C, NM_000157.3c.135G-S-T, NM_000157.3c.134G-S-C, NM_000157.3c.135G-S-T, NM_000157.3c.134G-S-C, NM_000157.3c.135G-S-T, NM_000157.3c.134G-S-C, NM_000157.3c.135G-S-T, NM_000157.3c.134G-S-C, NM_000157.3c.135C-S-T, NM_000157.3c.135c.914delC,	hepatosplenomegaly, anemia and thrombocytopenia, lung disease, and the absence of primary central nervous system disease. Of types 2 and 3 are characterized by the presence of primary neurologic disease. Type 2 has an onset before age two years, limited psychomotor development, and a rapidly progressive course with death by age two to four years. Type 3 may have onset before age two	600,25
			NM_000157.3c.586A>C, NM_000157.3c.5096>T, NM_000157.3c.508C>T, NM_000157.3c.508C>T, NM_000157.3c.487del(6, NM_000157.3c.481C>T, NM_000157.3c.476C>A, NM_000157.3c.475C>T, NM_000157.3c.475C>A, NM_000157.3c.475C>A, NM_000157.3c.647C>A, NM_000157.3c.647C>A, NM_000157.3c.65C>T, NM_000157.3c.259C>T, NM_000157.3c.254G>A, NM_000157.3c.647C>A, NM_000157.3c.647C>A, NM_000157.3c.647C>A, NM_000157.3c.647C>A, NM_000157.3c.647C>A, NM_000157.3c.84dupG	nore slowly progressive course, with survival into the third or fourth decade. The perinatal-lethal form is associated with ichthyosiform or collodion skin abnormalities or with nonimmune hydrops fetalis. The cardiovascular form is characterized by calcification of the aortic and mitral valves, mild splenomegally, corneal opacities, and supranuclear ophthalmoplegia. Cardiopulmonary complications have been described with all the clinical subtypes, although varying in frequency and severity. The incidence is 1/60,000 and the prevalence is approximately 1/100,000.	
GBE1	Glycogen storage disease, type 4	NM_000158.3	NM_000158.3:c:2052+1G>A, NM_000158.3:c:1883A>G, NM_000158.3:c:1774G>T, NM_000158.3:c:1804A>G, NM_000158.3:c:1570G>A, NM_000158.3:c:1570C>T, NM_000158.3:c:1543C>T, NM_000158.3:c:950C>C, NM_000158.3:c:771T>A, NM_000158.3:c:466_470delCGTAT	Clycogen storage disease, type 4 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the GEBI gene located on chromosomal region 3p12.2. The age of onset is infantile. This disease is characterized by failure to thrive; hepatomegaly, liver dysfunction, and progressive liver cirrhosis; hypotonia; cardiomyopathy and, finally, death. Clutaricaciduria, type 1	600,25
GCDH	Glutaricaciduria, type 1	NM_000159.3	NM_000159.3:c.74C>A, NM_000159.3:c.271+1G>A, NM_000159.3:c.383G>A, NM_000159.3:c.271+1G>A, NM_000159.3:c.542A>G, NM_000159.3:c.542A>G, NM_000159.3:c.542A>G, NM_000159.3:c.743C>T, NM_000159.3:c.542A>G, NM_000159.3:c.743C>T, NM_000159.3:c.751C>T, NM_000159.3:c.764C>T, NM_000159.3:c.751C>T, NM_000159.3:c.764C>T, NM_000159.3:c.764C>T, NM_000159.3:c.764C>T, NM_000159.3:c.764C>T, NM_000159.3:c.791C>T, NM_000159.3:c.1002.003641GA, NM_000159.3:c.1060G>A, NM_000159.3:c.1002.003641GA, NM_000159.3:c.1060G>A, NM_000159.3:c.1093G>A, NM_000159.3:c.1196G>C, NM_000159.3:c.1294C>T, NM_000159.3:c.1244-ZA>C, NM_000159.3:c.1244-ZA>C, NM_000159.3:c.1244-ZA>C, NM_000159.3:c.1244-ZA>C, NM_000159.3:c.1244-ZA>C, NM_000159.3:c.1246-ZA>C,	follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the GCDH gene located on chromosomal region 19p13.2. The age of onset is infantile or neonatal. This disease is characterized by encephalopathic crises resulting in striatal injury and a severe dystonic dyskinetic movement disorder. The prevalence is 1 in 100,000 births.	

GJB2	Deafness, autosomal recessive, type 1A	NM_004004.5	NM_004004.5:c.617A>G, NM_004004.5:c.551G>C, NM_004004.5:c.6517A>G, NM_004004.5:c.551G>A, NM_004004.5:c.45G5-A, NM_004004.5:c.45G5-A, NM_004004.5:c.45G5-A, NM_004004.5:c.45G5-A, NM_004004.5:c.45G5-A, NM_004004.5:c.45G5-A, NM_004004.5:c.45G5-A, NM_004004.5:c.45G5-A, NM_004004.5:c.35G5-AT, NM_004004.5:c.35G5-AT, NM_004004.5:c.35G5-AT, NM_004004.5:c.35G5-AT, NM_004004.5:c.310_323delAGAGAGTTCATCAAG, NM_004004.5:c.310_323delAGGAAGTTCATCAA, NM_004004.5:c.2593_300delAT, NM_004004.5:c.2590-AT, NM_004004.5:c.250G5-T, NM_004004.5:c.250G5-T, NM_004004.5:c.250G5-T, NM_004004.5:c.250G5-T, NM_004004.5:c.250G5-T, NM_004004.5:c.250G5-T, NM_004004.5:c.250G5-T, NM_004004.5:c.250G5-T, NM_004004.5:c.250G5-A, NM_004004.5:c.250G5-A, NM_004004.5:c.250G5-T, NM_004004.5:c.2	region Isql.2.11. The age or onset is infantile. This disease is characterized by congenital, non- progressive, mild-to- profound sensorineural hearing impairment. No other associated medical findings are present. Autosomal recessive	600,25
GJB6	Deafness, autosomal recessive, type 1B	NM_001110219.2	NM_001110219.2:c.485dupA, NM_001110219.2:c.443delC, NM_001110219.2:c.383_384delTA, NM_001110219.2:c.261dupA, NM_001110219.2:c.169C>T, NM_001110219.2:c.14C>T	region 13ql2.11. The age of onset is infantile. This disease is characterized by mild-to-profound sensorineural hearing impairment. No other associated medical findings are present.	600,25
GLB1	GM1-gangliosidosis, type 1	NM_001317040.1	NM_001317040.1c.1877A>G, NM_001317040.1c.1790C>T, NM_001317040.1c.1877A>G, NM_001317040.1c.1693G>T, NM_001317040.1c.1693G>T, NM_001317040.1c.1693G>T, NM_001317040.1c.1613G>T, NM_001317040.1c.1514G>A, NM_001317040.1c.1514G>A, NM_001317040.1c.1613G>T, NM_001317040.1c.1613G>T, NM_001317040.1c.1613G=T, NM_001317040.1c.1613B_1.319delCT, NM_001317040.1c.1613B_1.319delCT, NM_001317040.1c.1013B_1.319delCT, NM_001317040.1c.1013B_1.319delCT, NM_001317040.1c.1014SC>T, NM_001317040.1c.1013B_CST, NM_001317040.1c.1014SC>T, NM_001317040.1c.1014SC>T, NM_001317040.1c.74GG>A, NM_001317040.1c.74GG=A, NM_001317040.1c.7	In type 2 is late-infantile or juvenile and adult in type3. This disease is characterized by arrest/regression of neurological development, hypotonia, visceromegaly, macular cherry-red spots, dysostosis and coarse facial features. The prevalence is 1:100,000 a 200,000 newborn.	600,25
GLDC	Clycine encephalopathy	NM_000170.2	NM_000170.2:c.2405C>T, NM_000170.2:c.2284G>A, NM_000170.2:c.216G>A, NM_000170.2:c.1691C>T, NM_000170.2:c.1545G>C, NM_000170.2:c.1166C>T, NM_000170.2:c.322G>T	Glycine encephalopathy follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in CLDC gene located on chromosomal region 9p24.1. The age of onset is neonatal/infantile. This disease is characterized by lethargy or even coma, hypotonia, hiccups, myoclonic jerks, and breathing/swallowing disorders, with subsequent intellectual deficit, spasticity and intractable seizures. The prevalence is 11,000,000-91,000,000. Lethal congenital	600,25
GLEI	Lethal congenital contracture syndrome, type 1	NM_0010037221	NM_001003722.1:c.898-2A>G, NM_001003722.1:c.1412_1413delAG, NM_001003722.1:c.20511>C, NM_001003722.1:c.2069_2072delTTCT	contracture syndrome, type 1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the CLEI gene located on chromosomal region 9g34.11. The age of onset is neonatal. This disease is characterized by total fetal akinesia (detectable since the 13th week of gestation) accompanied by hydrops, micrognathia, pulmonary hypoplasia, pterygia and multiple joint contractures (usually flexion contractures in the elbows and extension in the knees), leading invariably to death before the 32nd week of gestation. Lack of anterior horn motoneurons, severe atrophy of the ventral spinal cord and severe skeletal muscle hypoplasia are characteristic	
GNE	Inclusion body myopathy, type 2 (Nonaka myopathy)	NM_001128227,2	NM_001128227.2:c.2228T>C, NM_001128227.2:c.2179G>A, NM_001128227.2:c.1891G>A, NM_001128227.2:c.1891G>A, NM_001128227.2:c.1820G>A, NM_001128227.2:c.0227-A, NM_001128227.2:c.830G>A, NM_001128227.2:c.830G>A, NM_001128227.2:c.478C>T	neuropathological findings, with no evidence of other organ structural anomalies. Inclusion body myopathy, type 2 (Nonaka myopathy) follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the GNE gene located on	600,25

GNPTAB	Mucolipidosis 2 alpha/beta	ı NM_024312.4	NM_024312.4:c.3663delG, NM_024312.4:c.3565C>T, NM_024312.4:c.35660.3561delAG, NM_024312.4:c.3503.3504delTC, NM_024312.4:c.3503.3504delTC, NM_024312.4:c.3503.3504delTC, NM_024312.4:c.3326dupA, NM_024312.4:c.31935c-9, NM_024312.4:c.2996delA, NM_024312.4:c.3503delA, NM_024312.4:c.1906dupA, NM_024312.4:c.1759C-T, NM_024312.4:c.196C-T, NM_024312.4:c.199delC, NM_024312.4:c.25C>T, NM_024312.4:c.10A>C	is caused by pathogenic variants in the GNPTAB gene located on chromosomal region 12q23.2. The age of onset is infantile. This disease is characterized by growth retardation, short stature, skeletal abnormalities, facial dysmorphism, stiff skin, developmental delay and cardiomegaly and that is lethal in childhood. The prevalence is 1:123,500-1625,500.	600,25
GPR179	Night blindness, congenital stationary (complete), type 1E, autosomal recessive	NM_001004334.3	NM_001004334.3:c.6847_6848delCT, NM_001004334.3:c.5693dupT, NM_001004334.3:c.5693_4700delAG, NM_001004334.3:c.4699_4700delAG, NM_001004334.3:c.323.3234delCT, NM_001004334.3:c.13824delCT, NM_001004334.3:c.1784+1C>A, NM_001004334.3:c.1368delT, NM_001004334.3:c.1236C>A, NM_001004334.3:c.984delC, NM_001004334.3:c.378delC	Congenital stationary night blindness type 1E follow an autosomal recessive pattern of inheritance and is caused by pathogenic variants in	600,25
CRM6	Night blindness, congenital stationary (complete), type 1B, autosomal recessive	NM_000843.3	NM_000843.3:c.2560C>T, NM_000843.3:c.2341G>A, NM_000843.3:c.2213_2219delCCAGAGG, NM_000843.3:c.2122C>T, NM_000843.3:c.1365C>T, NM_000843.3:c.1365C>T, NM_000843.3:c.1365C>T, NM_000843.3:c.136C>T, NM_000843.3:c.1214T>C, NM_000843.3:c.727dupG, NM_000843.3:c.712C>T	Congenital stationary night blindness type IB follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the GRMG gene located on chromosomal region 5q353. The age of onset is early infancy. This disease is characterized by hemeralopia with a moderate loss of visual	600,25
GUSB	Mucopolysaccharidosis, type 7	NM_000181.3	NM_000181.3:c.1881G>T, NM_000181.3:c.1856C>T, NM_000181.3:c.1831C>T NM_000181.3:c.1730G>T, NM_000181.3:c.1618G>T, NM_000181.3:c.1521G>A, NM_000181.3:c.1252G>A, NM_000181.3:c.1242+G>A, NM_000181.3:c.1242+G>A, NM_000181.3:c.1242+G>A, NM_000181.3:c.1244+G>A, NM_000181.3:c.1291.22016.c, NM_00181.3:c.1054-G>A, NM_000181.3:c.1291.22016.c, NM_00181.3:c.1055+G>T, NM_000181.3:c.1055+G>T, NM_00181.3:c.1052.856G>A, NM_000181.3:c.1055+G>T, NM_00181.3:c.1055+G>T, NM_00181.3:c.1055+G, NM_00181.3:c.1055+G, NM_00181.3:c.1055+G, NM_00181.3:c.1055+G, NM_00181.3:c	acuity. Mucopolysaccharidosis type 7 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the GUSB gene located on chromosomal region 7q112. The age of onset is variable. There are prenatal forms with non-immune hydrops fetalis, and severe neonatal forms with dysmorphism, hernias, hepatosplenomegaly, club feet, dysostosis, severe hypotonia and neurological disorders that ultimately lead to profound intellectual deficit and small stature in patients that survive. Finally, there are also very mild cases that are discovered during adolescence or adulthood following presentation with thoractic kyphosis. The prevalence is 1:250,000 in newborn. Isolated deficiency of long-losses.	
надна	LCHAD deficiency	NM_000182.4	NM_000182.4;c.2146+1G>A, NM_000182.4;c.2132dupC, NM_000182.4;c.1932.1794.delAT, NM_000182.4;c.17932.1794.delAT, NM_000182.4;c.16740=F, NM_000182.4;c.16740=F, NM_000182.4;c.1520+2_1620+5_delTAACG, NM_000182.4;c.1528G>C, NM_000182.4;c.1422dupT, NM_000182.4;c.132C>T, NM_000182.4;c.919-2A>G, NM_000182.4;c.274_278delTCATC	characterized in infancy/early childhood of hypoketotic hypoglycemia, metabolic acidosis, liver disease, hypotonia and, frequently, cardiac involvement with arrhythmias and/or cardiomyopathy. The	600,25
нвв	HBB-related hemoglobinopathy	NM_000518.4	NM_000518.4c: 110T>C, NM_000518.4c: 440_44idupAC, NM_000518.4c: 440A=T, NM_000518.4c: 440A=C, NM_000518.4c: 436A=T, NM_000518.4c: 436T>A, NM_000518.4c: 436T>A, NM_000518.4c: 436T>C, NM_000518.4c: 347T>C, NM_000518.4c: 346T>C, NM_000518.4c: 34	DNA variations in the HBB gene result in the production of different versions of beta-globin. Some of these variations may affect a person's health while other variations cause no noticeable signs or symptoms. Two of the most common HBB-related conditions are beta-thalassemia and sickle cell anemia (SCA). Beta thalassemia is caused by HBB gene mutations that prevent or decrease beta-globin production, subunits that make up hemoglobin. A lack of hemoglobin disrupts the normal development of red blood cells. A shortage of mature red blood cells can reduce the amount of oxygen that is delivered to tissues to	600,25

NM_0005I8.4:c.184A>1, NM_0005I8.4:c.1821>A, NM_0005I8.4:c.179A>C, satisfy the body's energy needs. A lack of oxygen in the body's tissues can lead to poor growth, organ NM_0005I8.4:c.133delC, NM_0005I8.4:c.127T>C, NM_0005I8.4:c.127T>C, NM_0005I8.4:c.127T>C, NM_0005I8.4:c.127T>C, NM_0005I8.4:c.127T>C, NM_0005I8.4:c.135delC, NM_0005I8.4:c.127T>C, NM_0005I8.4:c.137T>IBdelCC, NM_0005I8.4:c.1364-IMD_005IR.4:c.1364-IMD_005IR.4:c.1364-IMD_005IR.4:c.1364-IMD_005IR.4:c.1364-IMD_005IR.4:c.1364

pattern of inneritance and is caused by pathogenic variants in the HEXB gene located on chromosomal region 5q13.3. The age of onset is adult or infantile.

characterized by central nervous system degeneration, with startle reactions, early blindness, progressive motor and mental deterioration,

macrocephaly and cherryred spots on the macula The prevalence is 1/130.000. Alkaptonuria follows an

characterized by central 600.25

This disease is

HESX1	Growth hormone deficiency with pituitary anomalies	NM_003865.2	NM_003865.2:c.450_451delCA; NM_003865.2:c.445G>A; NM_003865.2:c.77T>C; NM_003865.2:c.18G>C	Growth hormone deficiency with pituitary anomalies follows an autosomal recessive pattern of inheritance and are caused by pathogenic variants in the HESXI gene located on chromosomal region 3p14.3. The age of onset is infantile. These diseases are characterized by short stature, cognitive alterations or delayed puberty. The incidence is 13,000 and 14,000 births.	
неха	Tay-Sachs disease	NM_000520.5	NM_000520.5:c.254-1G>C	Tay-Sachs disease follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the HEXA gene located on chromosomal region 15q25. The age of onset is infantile. There are three forms, type 1 (infantile), with a psychomotor retardation which is associated with hypotonia, amaurosis and megalencephaly. Type 2 is characterized by locomotor ataxia, behavioural disorders, and progressive loss of intellectual capacities. Type three (chronic form) shows spinocerebellar ataxia or spinal amyotrophy. The prevalence is 1 case per \$20000 live births.	600,25
неха	Tay-Sachs disease	NM_001318825.1	NM_0013188251:c.1570C-T, NM_0013188251:c.1561C-T, NM_0013188251:c.1544G-A, NM_0013188251:c.1543delC, NM_0013188251:c.1543delC, NM_0013188251:c.1543delT, NM_0013188251:c.1523delT, NM_0013188251:c.1528C-T, NM_0013188251:c.1528C-T, NM_0013188251:c.1457G-C, NM_0013188251:c.1470F-C, NM_0013188251:c.1293G-C, NM_0013188251:c.13013dupTATC, NM_0013188251:c.1293G-C, NM_0013188251:c.130140-TATC, NM_0013188251:c.13016-TATC, NM_0013188251:c.13016-TATC, NM_0013188251:c.13016-TATC, NM_0013188251:c.13016-TATC, NM_0013188251:c.13016-TATC, NM_0013188251:c.13016-TATC, NM_0013188251:c.13016-TATC, NM_0013188251:c.3016-CA, NM_001318825	Tay-Sachs disease follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the HEXA gene located on chromosomal region 15q23. The age of onset is infantile. There are three forms, type 1 (infantile), with a psychomotor retardation which is associated with hypotonia, amaurosis and megalencephaly. Type 2 is characterized by locomotor ataxia, behavioural disorders, and progressive loss of intellectual capacities. Type three (chronic form) shows spinocerebellar ataxia or spinal amyotrophy. The prevalence is 1 case per 320 000 live births. Sandhoff disease follows an autosomal recessive pattern of inheritance and	600,25

Sandhoff disease, infantile, NM_000521.3

NM_000187.3

iuvenile, and adult forms

HEXB

HGD

Alkaptonuria

Alkaptonuria follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the HGD gene located on chromosomal region 3ql3.33. The age of cast is infactly. NM_000187.3:c.1189-2A>G, NM_000187.3:c.1111dupC, onset is infantile. This NM_000187.3c.:1189-2-A>A, NM_000187.3c.:111dupC,

NM_000187.3c.:102A>C, NM_000187.3c.:111dupC,

NM_000187.3c.:102A>C, NM_000187.3c.:688C>T,

NM_000187.3c.:674G>A, NM_000187.3c.:688C>T,

NM_000187.3c.:674G>A, NM_000187.3c.:481C>A,

NM_00187.3c.:140C>T,

NM_00187.3c.:140C>T,

NM_00187.3c.:140C>T,

NM_00187.3c.:140C>T,

of the eye sclerae and the ear helix (ochronosis), and

NM_000521.3:c.115delG, NM_000521.3:c.171delG,
NM_000521.3:c.202_203insGG, NM_000521.3:c.298delC,
NM_000521.3:c.508c-T, NM_000521.3:c.7978-C, NM_000521.3:c.841C>T,
NM_000521.3:c.850c-T, NM_000521.3:c.1238_1.242delCAAAG,
NM_000521.3:c.1250C-T, NM_000521.3:c.1310_1.311delCA,
NM_000521.3:c.13545delT, NM_000521.3:c.1375G>T,

NM_000521.3:c.1345de11, NM_000521.3:c.1375d5 NM_000521.3:c.1380G5A, NM_000521.3:c.1517_1529dupCAAGTGCTGTTGG, NM_000521.3:c.1539_1540delCT

a disabling joint disease involving both the axial and peripheral joints (ochronotic arthropathy). The prevalence is 1:250,000-1:1.000.000 newborn. Mucopolysaccharidosis type 3C follows an autosomal recessive pattern of inheritance and pattern or innertrance and is caused by pathogenic variants in the HGSNAT gene located on chromosomal region 8p11.21. The age of onset is 600,25 infentile. This disease is NM_152419.2:c.493+1G>A, NM_152419.2:c.607C>T, NM_152419.2:c.848C>T, NM_152419.2:c.1030C>T, NM_152419.2:c.1250+1G>A, NM_152419.2:c.1378-1G>A, NM_152419.2:c.1530deIA, NM_152419.2:c.1553C>T, NM_152419.2:c.1622C>T HGSNAT Mucopolysaccharidosis type 3C (Sanfilippo C) NM_152419.2 infantile. This disease is characterized by defective characterized by defective or missing enzymes to break down mucopolysaccharides are missing or are defective. The prevalence is <1:70.000 newborn. Tyrosinemia type 3 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the HPD gene variants in the HPD gene located on chromosomal region 12q24.31. The age of 600,25 onset is infantile. This disease is characterized by intellectual deficit and NM 002150 2:c 987delA NM 002150 2:c 774T>G HPD Tyrosinemia, type 3 NM_002150.2 NM_002150.2:c.600C>G ataxia. The prevalence is 1:100,000-1:120,000 newborn. Charcot-Marie-Tooth disease, axonal, type 2S follows an autosomal follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the IGHMBP2 gene located on chromosomal region 11q13.3. The age of parts to an abolishmen. NM_002180.2:c.12iC>T, NM_002180.2:c.638A>G, NM_002180.2:c.66idelA, dult or adolescent. This disease is characterized by NM_002180.2:c.112iC>T, NM_002180.2:c.638A>G, NM_002180.2:c.66idelA, progressive distal muscle NM_002180.2:c.1540G>A, NM_002180.2:c.1738G>A, weakness and atrophy of both the lower and upper limbs—st IGHMBP2 Charcot-Marie-Tooth disease, axonal, type 2S NM_002180.2 limbs, absent or reduced deep tendon reflexes, mild sensory loss, foot drop, and pes cavus leading eventually to wheelchair dependence. Some patients present with early hypotonia and delayed motor development. Scoliosis and variable autonomic disturbances may be associated. The prevalence is below 1/1,000,000. Joubert syndrome type 1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the INPPSE gene located on chromosomal region 934-3. The age of onset is early infantile. This disease is characterized to accept the pattern of the pathogene in the pattern of the pathogene in the pattern of the pathogene in the pathogene is characterized congenital malformation of the malformation of the brainstem and agenesis of the cerebellar vermis (molar tooth sign) leading to an abnormal respiratory 600,25 pattern, nystagmus, NM_019892.5:c.1879C>T, NM_019892.5:c.1688G>A, NM_019892.5:c.1543C>T, NM_019892.5:c.1304G>A, NM_019892.5:c.1132C>T, NM_019892.5:c.855_856insCG INPPSE Joubert syndrome, type 1 NM_019892.5 hypotonia, mental nypotonia, mental retardation, ataxia, and delay in achieving motor milestones. Other variable features include retinal dystrophy (manifesting with either Leber congenital amaurosis or dystrophy) and nephronophthisis (usually juvenile). The prevalence is 1:100,000. Diabetes mellitus, insulinresistant, with acanthosis nigricans type A follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the INSR gene located on chromosomal region 19p13.2. The age of onset is infantile. This disease is characterized by the triad of hyperinsulinemia, acanthosis nigricans (skin lesions associated with insulin resistance), and signs of hyperandrogenism in females without lipodystrophy and who are variants in the INSR gene lipodystrophy and who are ilpodystropny and who a not overweight. It is generally diagnosed in young women with marked signs of hyperandrogenism, but insulin resistance and Diabetes mellitus, insulin-resistant, with acanthosis NM_000208.3 nigricans, type A NM_000208.3:c.3680G>C, NM_000208.3:c.3079C>T, NM_000208.3:c.2668C>T, NM_000208.3:c.1114C>T, NM_000208.3:c.172G>A INSP 600,25 insulin resistance and acanthosis nigricans may be observed in men and in childhood. Acromegaloid facies or muscular cramps are sometimes associated. Hyperinsulinemia, a highorical marker for biological marker for insulin resistance, is often

a disabling joint disease

associated with glucose tolerance defects over the

	Epidermolysis bullosa,		NM_000213.4:c.112T>C, NM_000213.4:c.182G>A, NM_000213.4:c.1150delG, NM_000213.4:c.1660C>T, NM_000213.4:c.1664T>C, NM_000213.4:c.2608delC, NM_000213.4:c.27972-G, NM_000213.4:c.3371.3331delACTGCACCGCA	course of the disease, and diabetes progressively sets in. Hyperandrogenism (associated with polycystic ovarian syndrome (see this term) or ovarian hyperthecoses) leads to fertility problems. The prevalence is <11,000,000. Junctional epidermolysis bullosa with pyloric atresia follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ITGB4 gene located on chromosomal region 17q251. The age of onset is	
ITGB4	junctional, with pyloric atresia	NM_000213.4	NM_000213.4:c.2792G>A, NM_000213.4:c.3321_3331delACTGGACCGGA, NM_000213.4:c.3574G>A, NM_000213.4:c.3793+1GA, NM_000213.4:c.3801dupT, NM_000213.4:c.3894C>T, NM_000213.4:c.4823GCA, NM_000213.4:c.4823GC>T, NM_000213.4:c.4823GC>T, NM_000213.4:c.5329+2T>C	early infantile. This disease is characterized by generalized blistering at birth and congenital atresia of the pylorus and rarely of other portions of the gastrointestinal tract. More than 100 cases have been reported around the world.	
IVD	Isovaleric acidemia	NM_002225.3	NM_002225.3c.2T>G, NM_002225.3c.134T>C, NM_002225.3c.157C>T, NM_002225.3c.158C>A, NM_002225.3c.158G>C, NM_002225.3c.158G>C, NM_002225.3c.158G>C, NM_002225.3c.243-16-A, NM_002225.3c.36G>C, NM_002225.3c.243G-A, NM_002225.3c.406.407delTG, NM_002225.3c.454_4.437dupATGA, NM_002225.3c.507delG, NM_002225.3c.507delG, NM_002225.3c.507delG, NM_002225.3c.507delG, NM_002225.3c.507delG, NM_002225.3c.507delG, NM_002225.3c.507delG, NM_002225.3c.6507delG, NM_002225.3c.6507delG, NM_002225.3c.6507delG, NM_002225.3c.6507delG, NM_002225.3c.627delT, NM_002225.3c.6507delG, NM_002225.3c.6507delG, NM_002225.3c.627delT, NM_002225.3c.697delG, NM_002225.3c.1145T-C, NM_002225.3c.1145T-4delTTGGTGA, NM_002225.3c.1183C>T, NM_002225.3c.1188delT, NM_002225.3c.1192C>T, NM_002225.3c.1208A>G	Isovaleric academia follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the IVD gene located on chromosomal region 15q15.1. The age of onset is neonatal. This disease is characterized by vomiting, dehydration, coma and abnormal movements. The prevalence is 1/100,000. Severe combined	600,25
Ј АКЗ	Severe Combined Immunodeficiency, autosomal recessive, T- negative/B-positive type	NM_000215.3	NM_000215.3:c.1837C>T, NM_000215.3:c.1765G>A, NM_000215.3:c.1835C>A, NM_000215.3:c.1333C>T, NM_000215.3:c.1172_1173insG, NM_000215.3:c.299A>G	immunodeficiency, T-B+ type follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the JAK3 gene located on chromosomal region 19p13.11. The age of onset is	600,25
KCNJI	Bartter syndrome, type 2	NM_000220.4	NM_000220.4:c:1014delA, NM_000220.4:c:1012C>T, NM_000220.4:c:996_999delAAAG, NM_000220.4:c:942T>G, NM_000220.4:c:597C>G, NM_000220.4:c:500G>T, NM_000220.4:c:592G>A, NM_000220.4:c:500G>A, NM_000220.4:c:372T>A, NM_000220.4:c:322G>C, NM_000220.4:c:237C>G	pathogens. The incidence is 1/100,000 and 1/1,000,000. Bartter syndrome, type 2 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the KCNJI gene located on chromosomal region 11q24.3. The age of onset is antenatal. This disease is	600,25
KCNV2	Retinal cone dystrophy, type 3B	NM_133497.3	NM_133497.3:c 226C>T, NM_133497.3:c.325C>T, NM_133497.3:c.357dupC, NM_133497.3:c.42C>T, NM_133497.3:c.491T>C, NM_133497.3:c.647C>T, NM_133497.3:c.961T>C, NM_133497.3:c.1016_1024delACCTGGTGG, NM_133497.3:c.1133dupT, NM_133497.3:c.1376G>A	Retinal cone dystrophy, type 3B follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the KCNV2 gene located on 9p24.2. The age of onset is in the first or second decade of life. This disease is characterized by is characterized by onset in the first or second decade of life of very marked photophobia, myopia, reduced color vision along the red-green axis with relatively preserved tritan discrimination, and central discrimination, and central secotomats with peripheral soctomats with peripheral	600,25
				widespread sensitivity loss predominating in the superior visual field. Nyctalopia is a later feature of the disorder. There is often retiral pigment epithelium disturbance at the macula with a normal retinal periphery. LAMA2-related muscular dystrophy 1A follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the LAMA2 gene located on chromosomal region 6q22.33. LAMA2-related muscular dystrophy is a disorder that causes weakness and atrophy of skeletal muscles. This condition varies in severity, from a severe, early-onset type to a milder, late-onset type to a milder, late-onset dystrophy is apparent at birth or within the first few months of life, called	

NM_000426.3:c.112+1G>A, NM_000426.3:c.184G>T, NM_000426.3:c.185delC, NM_000426.3:c.1050delT, NM_000426.3:c.1050delT, NM_000426.3:c.1050delT, NM_000426.3:c.1050delT, NM_000426.3:c.1050delT, NM_000426.3:c.1050delT, NM_000426.3:c.2190delT, NM_000426.3:c.2750-C, NM_000426.3:c.250delT, NM_000426.3:c.250delT, NM_000426.3:c.250delT, NM_000426.3:c.257G>A, NM_000426.3:c.257G>A, NM_000426.3:c.257G>A, NM_000426.3:c.357G>T, NM_000426.3:c.757GelCT, NM_00426.3:c.757GelCT, NM_000426.3:c.757GelCT, NM_000426.3:c.757GelCT, NM_000426.3:c.757GelCT, NM_000426.3:c.757GelCT, NM_000426.3:c.757GelCT, NM_000426.3:c.757GelCT, NM_000426.3:c.757GelCT, NM_000426.3:c.757GelT, NM_00426.3:c.757GelT, NM_000426.3:c.757GelT, NM_00426.3:c.757GelT, NM_00426.3:c.757GelT, NM_00426.3:c.757GelT, NM_00426.3:c.757GelT, NM_00426.3:c.757GelT, NM_00426.3:c.757GelT, NM_00426.3:c.757GelT, NM_00426.3:c.757GelT, NM_00426.3:c.7 NM_000426.3:c.112+1G>A, NM_000426.3:c.184G>T,

LAMA2-related muscular LAMA2 dystrophy

NM 000426.3

Junctional epidermolysis LAMB3 bullosa, Herlitz and non-Herlitz type

NM_000228.2

NM_000228.2:c.3228+1G>T, NM_000228.2:c.3228+1G>A. NM_000228.2c.3228+1G-T, NM_000228.2c.3228+1G-A, NM_000228.2c.3286+1C, NM_000228.2c.3286+1C, NM_000228.2c.3286+1AG, NM_000228.2c.1383G-A, NM_000228.2c.387.3588delAG, NM_000228.2c.357.3542delT, NM_000228.2c.357delT, NM_000228.2c.357delT, NM_000228.2c.357delT, NM_000228.2c.357delT, NM_000228.2c.3587delT, NM_000288.2c.3587delT, NM_000288.2c.3587delT, NM_000288.2c.3587delT, NM_000288.2c.3587delT, NM_000288.2c.3587delT, NM_000288.2c.3587delT, NM_000288.2c.3587delT, NM_000288.2c.3587d NM_000228.2:c.124C>T

congenital muscular dystrophy type 1A (607855). Patients show hypotonia, poor suck and cry, and delayed motor development; most never achieve independent ambulation. Most patients also have periventricular white matter abnormalities on brain imaging, but mental imaging, but mental retardation and/or seizures 600,25 occur only rarely.

Symptoms of late-onset LAMA2-related muscular dystrophy become evident later in childhood or later in childhood or adulthood, and are similar to those of a group of muscle disorders classified as autosomal recessive limb-girdle muscular dystrophies, type 23. This group is characterized by slowly progressive proximal muscle proximal muscle weakness primarily affecting the lower lim and resulting in gait difficulties. Additional features include white limbs matter abnormalities on brain imaging, increased serum creatine kinase, and dystrophic features with partial LAMA2 deficiency on muscle biopsy. Some patients may have additional neurologic have additional neurologic features, including executive deficits, seizures, and peripheral neuropathy. Patients remain ambulatory well into adulthood. The prevalence is 1/30,000. Junctional epidermolysis bullosa follows an autosomal recessive pattern of inheritance and pattern of inheritance and is caused by pathogenic variants in the LAMB3 gene located on chromosomal region lq32.2. The age of onset is neonatal/infancy. Junctional epidermolysis bullosa (JEB) is a group of genetic conditions that cause the skin to be very fragile and to blister easily. Blisters and skin erosions form in response to minor form in response to mino injury or friction, such as rubbing or scratching. Researchers classify junctional epidermolysis bullosa into two main types based on severity: 600,25 Herlitz JEB and non-Herlitz JEB and non-Herlitz JEB. Herlitz type is more severe phenotype characterized by blisters and erosions, localized to the skin and mucous membranes and often results in early death. More results in early death. More than 80 mutations in the LAMB3 gene have been identified in people with Herlitz JEB. Other LAMB3 gene mutations cause the milder form non-Herlitz JEB, disease characterized JEB, disease characterized by generalized skin blistering, atrophic scarring, nail dystrophy of nail absence, and enamel hypoplasia, with extracutaneous involvement. LMNA-related disorders, autosomal recessive, are caused by pathogenic variants in the LMNA gene variants in the LMNA gen located on chromosomal region 1q22, and include Charcot-Marie-Tooth disease, type 2B1, Emery-Dreifuss muscular dystrophy type 3, mandibuloacral dysplasia, mandibuloacral dysplasia, lethal restrictive dermopathy among others. Charcot-Marie-Tooth disease constitutes a clinically and genetically a climically and genetically heterogeneous group of hereditary motor and sensory neuropathies. Emery-Dreifuss muscular dystrophy is characterized classically by the triad of weakness of the shoulder weakness of the shoulder and pelvic girdle muscles, contractures of the elbows, neck, and Achilles tendon, and cardiac involvement, most commonly arrhythmias. Mandibuloacral dysplasia 600,25 Mandibuloacral dysplasia is characterized by growth retardation, craniofacial anomalies with mandibular hypoplasia, skeletal abnormalities with progressive osteolysis of the distal phalanos.

of the distal phalanges and clavicles, and pigmentary skin changes Restrictive dermopathy is

LMNA-related disorders.

LMNA

autosomal recessive

NM_001282626.1 NM_001282626.1;c.1818+6C>T

LMNA	LMNA-related disorders, autosomal recessive	NM_170707.3	NM_170707.3:c.4191>C, NM_170707.3:c.1072G>A, NM_170707.3:c.1228C>T, NM_170707.3:c.1366a>C, NM_170707.3:c.1410>T, NM_170707.3:c.1488+1G>A, NM_170707.3:c.1583C>A, NM_170707.3:c.1580C>A, NM_170707.3:c.1583C>A, NM_170707.3:c.1626G>C	a rare, lethal genodermatosis characterized by thin, tightly adherent translucent skin with erosions at flexure sites, superficial vessels, typical facial dysmorphism, and generalized joint ankylosis. LMNA-related disorders, autosomal recessive, are caused by pathogenic variants in the LMNA gene located on chromosomal region 1q22, and include charcot-Marie-Tooth disease, type 2B1, Emery-Dreifuss muscular dystrophy type 3, mandibuloacral dysplasia, lethal restrictive dermopathy among others. Charcot-Marie-Tooth disease constitutes a clinically and genetically heterogeneous group of hereditary motor and sensory neuropathies. Emery-Dreifuss muscular dystrophy is characterized classically by the triad of weakness of the shoulder and pelvic girdle muscles, contractures of the elbows, neck, and Achilles tendon, and cardiac involvement, most commonly arrhythmias. Mandibuloacral dysplasia is characterized by growth retardation, cranifacial anomalies with mandibular hypoplasia, skeletal abnormalities with mandibular hypoplasia, skeletal abnormalities with mandibular stellar progressive osteolysis of the distal phalanges and clavicles, and pigmentary skin changes. Restrictive dermopathy is a rare, lethal genodermatosis characterized by thin, tightly adherent translucent skin with erosions at flexure sites, superficial vessels, typical facial dysmorphism, and generalized joint ankylosis. Osteoporosis-pseudoglioma syndrome	600,25
LRP5	Osteoporosis- pseudoglioma syndrome	NM_002335.3	NM_002335.3:c.804_813delGGGGAAGAGG, NM_002335.3:c.1453G>T, NM_002335.3:c.1486delG, NM_002335.3:c.1481G>A, NM_002335.3:c.1708G>T, NM_002335.3:c.1708G>T, NM_002335.3:c.2026-A, NM_002335.3:c.254C>G, NM_002335.3:c.2557C>T, NM_002335.3:c.4651G>A	follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the LRPS gene located on chromosomal region 10 ₄ 13.2. The age of onset is infantile. This disease is characterized by congenital or infancyonset blindness and severe juvenile-onset osteoporosis and spontaneous fractures. The prevalence is	600,25
MANZBI	Mannosidosis, alpha-, types I and II	NM_000528.3	NM_000528.3:c.2686_2687delCTinsG, NM_000528.3:c.2436+2T>C, NM_000528.3:c.2426+2T>C, NM_000528.3:c.2398G>A, NM_000528.3:c.2596C>T, NM_000528.3:c.2278C>T, NM_000528.3:c.219C>T, NM_000528.3:c.2013delT, NM_000528.3:c.1915C>T, NM_000528.3:c.193C>T, NM_000528.3:c.193C>T, NM_000528.3:c.1830+1G>C, NM_00528.3:c.1780C>T, NM_000528.3:c.384G>A, NM_000528.3:c.1A>G	1:2,000,000. Alpha-mannosidosis follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the MAN2BI gene located on chromosomal region 19p13.2. The age of onset is infantile. This disease is characterized by immunodeficiency, facial and skeletal abnormalities, hearing impairment and intellectual disability. The prevalence is 1:1,000,000-9:1,000,000. 3-methylcrotonyl-CoA carboxylase deficiency	
MCCC2	3-Methylcrotonyl-CoA carboxylase type 2, deficiency	NM_022132.4	NM_022132.4:c.295G>C, NM_022132.4:c.380C>G, NM_022132.4:c.464G>A, NM_022132.4:c.499T>C, NM_022132.4:c.517dupf, NM_022132.4:c.634GelG, NM_022132.4:c.537dupf, NM_022132.4:c.838G>T, NM_022132.4:c.939C>C, NM_022132.4:c.994C>T, NM_022132.4:c.1015G>A, NM_022132.4:c.1065A>T, NM_022132.4:c.10724IG>A, NM_022132.4:c.1577dupf, NM_022132.4:c.1580G>A	type 2 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the MCCC2 gene located on chromosomal region 5q13.2. The age of onset is neonatal. This disease is characterized by a highly variable clinical picture ranging from neonatal onset with severe neurological involvement to asymptomatic adults.	
MED25	Basel-Vanagait-Smirin- Yosef syndrome	NM_030973.3	NM_030973.3:c.320delG, NM_030973.3:c.1366C>T	The prevalence is 1:75,000 newborn. Basel-Vanagait-Smirin-Yosef syndrome follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the MED25 gene located on chromosomal region 19q13.33. The age of onset neonatal/infantile. This syndrome is characterized by eye, brain, cardiac and palatal abnormalities as well as growth retardation, microcephaly and severe intellectual disability. Familial Mediterranean fever follows an autosomal recessive pattern of inheritance and is caused	600,25

MEFV	Familial Mediterranean fever, AR	NM_000243.2	NM_000243.2:c.2282G>A, NM_000243.2:c.2230G>T, NM_000243.2:c.2730G>T, NM_000243.2:c.27717>C, NM_000243.2:c.2084A>G, NM_000243.2:c.2806A>G, NM_000243.2:c.2806A>G, NM_000243.2:c.2806A>G, NM_000243.2:c.2940G>C, NM_000243.2:c.2940G>C, NM_000243.2:c.2940G>C, NM_000243.2:c.187C>G, NM_000243.2:c.187C>G, NM_000243.2:c.187C>G, NM_000243.2:c.187C>G, NM_000243.2:c.501G>C, NM_000243.2:c.163dupA	by patriogenic variants in the MEFV gene located on chromosomal region 16p13.3. The age of onset is infantile or adult (before the age of 30). This disease is characterized by recurrent short episodes of fever and serositis resulting in pain in the abdomen, chest, joints and muscles. The prevalence is 110,000-	
MERTK	Retinitis pigmentosa type 38	NM_006343.2	NM_006343.2:c.1605-2A>G, NM_006343.2:c.2070_2074delAGGAC, NM_006343.2:c.2189+1G>T, NM_006343.2:c.271_2214delCTGT, NM_006343.2:c.2323C>T, NM_006343.2:c.2785_2786dupTA	\$10,000. Retinitis pigmentosa type 38 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the MERTK gene located on chromosomal region 23. The age of onset is infantile. This disease is characterized by. This disease is characterized by night blindness, followed by a progressive loss of peripheral vision in the daylight period and leading to blindness.	600,25
MFRP	Microphthalmia, isolated type 5	NM_031433.3	NM_031433.3:c:1149dupC, NM_031433.3:c:1124+1G>T, NM_031433.3:c:545T>C, NM_031433.3:c:523C>T, NM_031433.3:c:498delC	Microphthalmia, isolated type 5 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the MFRP gene located on chromosomal	600,25
MKKS	Bardet-Biedl syndrome type 6	NM_018848.3	NM_018848.3:c.1436C>G, NM_018848.3:c.1225_1226delGG, NM_018848.3:c.830T>C, NM_018848.3:c.353delG	type 6 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the MKKS gene located on chromosomal region 20p12.2 The age of onset is antenatal or infacy. This disease is characterized by a combination of clinical signs: obesity, pigmentary retinopathy, post-axial polydactyly, polycystic kidneys, hypogenitalism, and learning disabilities, many of which appear several years after disease onset. Clinical expression is variable but most	600,25
MKS1	Bardet-Biedl syndrome type 13	NM_001321269.1	NM_001321269:1:c:1024+1G>A, NM_001321269:1:c:508C>T	patients manifest the majority of clinical signs during the disease course. Bardet-Biedl syndrome type 13 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the MKSI gene located on chromosomal region 1702.2 The age of onset is antenatal or infacy. This disease is characterized by a combination of clinical signs: obesity, pigmentary retinopathy, post-axial polydactlyly, polycystic kidneys, hypogenitalism, and learning disabilities, many of which appear several years after disease onset. Clinical expression is variable but most patients manifest the majority of clinical signs during the disease course.	600,25
ммаснс	Methylmalonic aciduria and homocystinuria, cbIC type	NM_015506.2	NM_015506.2:c.27IdupA, NM_015506.2:c.331C>T, NM_015506.2:c.347T>C, NM_015506.2:c.384.390delTAC, NM_015506.2:c.394C>T, NM_015506.2:c.3475>C, NM_015506.2:c.496C>T, NM_015506.2:c.496C>T, NM_015506.2:c.496C>T, NM_015506.2:c.547_548.delGT, NM_015506.2:c.608C>A, NM_015506.2:c.609C>A, NM_015506.2:c.609C>A, NM_015506.2:c.609C>A, NM_015506.2:c.616C>T, NM_015506.2:c.616C>T, NM_015506.2:c.616C>T, NM_015506.2:c.616C>T, NM_015506.2:c.616C>T, NM_015506.2:c.658_660delAAG, NM_015506.2:c.688C>T	Vitamin B12-responsive methylmalonic acidemia type cb1 B follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the MMAB gene located on chromosomal region 12q24-31. The age of onset is early infantile. This disease is characterized by developmentally delayed with other features that include hypotonia, seizures, hypoglycaemia, metabolic acidosis, cardiomyopathy and	
MOCS2	Molybdenum cofactor deficiency B	NM_004531.4	NM_004531.4:c.567A>C, NM_004531.4:c.539_540delAA, NM_004531.4:c.502G>A, NM_004531.4:c.377+1C>A, NM_004531.4:c.106_107delAT, NM_004531.4:c.58delT, NM_004531.4:c.3G>A	diarrhoea. The prevalence is <13,000,000. Molybdenum cofactor deficiency type B follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the MOCS2 gene located on chromosomal region 5q11.2. This disease is characterized by severe neurological abnormalities, dislocated ocular early death. Molybdenum cofactor deficiency type B follows	600,25
MOCS2	Molybdenum cofactor	NM 176806.3	NM 176806.3:c.16C>T	an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the MOCS2 dene located on	600.25

	аетісіепсу в	=	-	chromosomal region	
				5q11.2. This disease is characterized by severe neurological	
				abnormalities, dislocated ocular early death.	
				Abetalipoproteinemia follows an autosomal	
				recessive pattern of inheritance and is caused by pathogenic variants in	
				the MTTP gene located on chromosomal region 4q23.	
MTTP	Abetalipoproteinemia	NM_001300785.1	NM_001300785.1:c.789_790delCA, NM_001300785.1:c.1700G>A, NM_001300785.1:c.1850G>T, NM_001300785.1:c.1948+1G>A, NM_001300785.1:c.2102delC, NM_001300785.1:c.2674G>T	The age of onset is infantile. This disease is	600,25
			MI_00/300703.i.e.21/20010, MII_00/300703.i.e.20740-1	characterized by growth delay, malabsorption,	
				hepatomegaly, and neurological and neuromuscular	
				manifestations. The prevalence is <1:1,000,000.	
				Methylmalonic acidemia follows an autosomal	
			NM_000255.3:c.2150G>T, NM_000255.3:c.2080C>T, NM_000255.3:c.1924G>C, NM_000255.3:c.1871A>G,	recessive pattern of inheritance and is caused by pathogenic variants in	
			NM_000255.3:c.1867G>A, NM_000255.3:c.1741C>T, NM_000255.3:c.1658delT, NM_000255.3:c.1445-2A>G,	the MUT gene located on chromosomal region	
MUT	Methylmalonic aciduria, mut(0) type	NM_000255.3	NM_000255.3:c.1420C>T, NM_000255.3:c.1399C>T, NM_000255.3:c.1280G>A, NM_000255.3:c.1207C>T, NM_000255.3:c.1181T>A, NM_000255.3:c.1106G>A,	6p12.3. The age of onset is very early infantile. This	600,25
			NM_000255.3:c.914T>C, NM_000255.3:c.682C>T, NM_000255.3:c.671_678dupAATTTATG, NM_000255.3:c.655A>T,	disease is characterized by recurrent ketoacidotic comas or transient	
			NM_000255.3:c.643C>A, NM_000255.3:c.607G>A, NM_000255.3:c.572C>A, NM_000255.3:c.313T>C, NM_000255.3:c.280G>A, NM_000255.3:c.278G>A, NM_000255.3:c.91C>T	vomiting, dehydration, hypotonia and intellectual	
			NW_UUU235.3.C.260G / A, NW_UUU235.3.C.276G / A, NW_UUU235.3.C.51C / 1	deficit, which does not respond to administration	
				of vitamin B12. Mevalonic aciduria follows an autosomal recessive	
				pattern of inheritance and is caused by pathogenic	
				variants in the MVK gene located on chromosomal	
MVK	Mevalonic aciduria	NM_000431.3	NM_000431.3:c.59A>C, NM_000431.3:c.185G>A, NM_000431.3:c.494C>T, NM_000431.3:c.803T>C, NM_000431.3:c.902A>C,	region 12q24.11. The age of onset is infantile. This disease is characterized by	600.35
IVIV	Wevaloriic aciduria	1414_000451.5	NM_000431.3:c.928G>A, NM_000431.3:c.1000G>A, NM_000431.3:c.1129G>A	psychomotor retardation, failure to thrive,	000,23
				progressive cerebellar ataxia, dysmorphic	
				features, progressive visual impairment and recurrent febrile crises. The	
				prevalence is <1:1,000,000. Deafness autosomal	
				recessive type 3 follows an autosomal recessive	
			NM_016239.3:c.625G>T, NM_016239.3:c.755dupA, NM_016239.3:c.3313G>T, NM_016239.3:c.3336delG,	pattern of inheritance and is caused by pathogenic variants in the MYO15A	
	Deafness, autosomal		NM_016239.3:c.3385C>T, NM_016239.3:c.3693-2A>G, NM_016239.3:c.3756+1G>T, NM_016239.3:c.4751_4752dupTC,	gene located on chromosomal region	
MYO15A	recessive type 3	NM_016239.3	NM_016239.3:c.5326C>T, NM_016239.3:c.5492G>T, NM_016239.3:c.6004delG, NM_016239.3:c.6864_6874delGGACCTGGAGC, NM_016239.3:c.8148G>T,	17pl1.2. The age of onset is infantile, etc/. This disease is characterized by mild-	600,25
			NM_016239.3:c.8410A>T, NM_016239.3:c.8548C>T, NM_016239.3:c.958_9961delGACT, NM_016239.3:c.10573delA	to-profound sensorineural hearing impairment with	
				no associated visible abnormalities of the	
				external ear or any related medical problems. Deafness autosomal	
				recessive type 30 follows an autosomal recessive	
				pattern of inheritance and is caused by pathogenic	
			NM_017433.4:c.1A>G, NM_017433.4:c.732-2A>G, NM_017433.4:c.770C>G, NM_017433.4:c.1086T>G, NM_017433.4:c.1193C>A, NM_017433.4:c.1777-	variants in the MYO3A gene located on chromosomal region	
MYO3A	Deafness, autosomal recessive type 30	NM_017433.4	12C>A, NM_017433.4:c.1953delC, NM_017433.4:c.2243delA, NM_017433.4:c.2506-1G>A, NM_017433.4:c.2793+2T>A, NM_017433.4:c.3154C>T,	10p12.1. The age of onset is infantile. This disease is	600,25
			NM_017433.4:c.4586+2T>G, NM_017433.4:c.4730+1G>A	characterized by mild-to- profound sensorineural	
				hearing impairment with no associated visible abnormalities of the	
				external ear or any related medical problems.	
				Deafness autosomal recessive type 37 follows	
				an autosomal recessive pattern of inheritance and is caused by pathogenic	
				variants in the MYO6 gene located on chromosomal	
MYO6	Deafness, autosomal recessive type 37	NM_004999.3	NM_004999.3:c.1452dupT, NM_004999.3:c.2907_2909delAGA, NM_004999.3:c.3496C>T, NM_004999.3:c.3808C>T	region 6q14.1. The age of onset is infantile. This disease is characterized by	600,25
				mild-to-profound sensorineural hearing	
				impairment with no associated visible abnormalities of the	
				external ear or any related medical problems.	
				Usher syndrome type 1B follows an autosomal	
			NM_000260.3:c.3G>A, NM_000260.3:c.133-2A>G, NM_000260.3:c.448C>T, NM_000260.3:c.494C>T, NM_000260.3:c.634C>T, NM_000260.3:c.635G>A,	recessive pattern of inheritance and is caused by pathogenic variants in	
			NM_000260.3:c.640G>A, NM_000260.3:c.731G>C, NM_000260.3:c.1184G>A, NM_000260.3:c.1344-1G>A,	the MYO7A gene located on chromosomal region	
МУО7А	Usher syndrome, type 1B	NM_000260.3	NM_000260.3:c.1797G>A, NM_000260.3:c.1884C>A, NM_000260.3:c.1996C>T, NM_000260.3:c.2476G>A,	11q13.5. The age of onset is	600,25
			NM_000260.3:c:3504-1G>c, NM_000260.3:c:3508G>A, NM_000260.3:c:3596dupT, NM_000260.3:c:3719G>A, NM_000260.3:c:3764delA, NM_000260.3:c:4024delT,	congenital, bilateral, profound sensorineural	
			NM_000260.3:c.5392C>T, NM_000260.3:c.5618G>A, NM_000260.3:c.5824G>T, NM_000260.3:c.5886_5889delCTTT,	hearing loss, vestibular areflexia, and adolescent-	
			NM_000260.3:c.5967C>G, NM_000260.3:c.6025delG	onset retinitis pigmentosa. The prevalence is 1:100,000-9:100,000.	
				Schindler disease follows an autosomal recessive	
				pattern of inheritance and is caused by pathogenic	
				variants in the NAGA gene located on chromosomal region 22q13.2. The age of	
				J. o LEGIO. E. THE age Of	

NAGA	Schindler disease, type I	NM_000262.2	NM_000262.2:c.986G>A, NM_000262.2:c.985C>T, NM_000262.2:c.973G>A, NM_000262.2:c.577G>T	onset is infantile. This disease is characterized by early-onset neuroaxonal dystrophy and neurological signs (convulsion during fever, epilepsy, psychomotor retardation and hypotonia). NAGA deficiency is typically classified in three main phenotypes: NAGA deficiency type I (Schindler disease type I) with severe manifestations; NAGA deficiency type II (Kanzazi disease or Schindler disease type II) which is mild: NAGA deficiency type III (Schindler disease type III) which is mild: NAGA deficiency type III (Schindler disease type III) chindler disease type III) chindler disease type III haracterized by mild-to-moderate neurologic manifestations. NAGA deficiency results in the increased urinary excretion of glycopeptides and oligosaccharides containing alpha-N-acetylogalactosaminyl	600,25
NEB	Nemaline myopathy type 2, autosomal recessive	NM_001271208.1	NM_001271208.1:c.12238_12239delAT, NM_001271208.1:c.8031_8041delAAATAAACGAG, NM_001271208.1:c.61054dupT, NM_001271208.1:c.2173G>T, NM_001271208.1:c.843T>G	moleties. Nemaline myopathy type 2 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the NEB gene located on chromosomal region 2q23.3. The age of onset is infantile or adult. This disease is characterized by hypotonia, weakness and depressed or absent deep tendon reflexes, with pathologic evidence of nemaline bodies (rods) on muscle biopsy. The prevalence is 1:100,000- 9:100,000 and the incidence is 1/50.000 newborn.	600,25
NMNATI	Leber congenital amaurosis type 9	NM_001297778.1	NM_001297778.1:c.25G>A, NM_001297778.1:c.451G>T, NM_001297778.1:c.57G>A, NM_001297778.1:c.507G>A, NM_001297778.1:c.710G>T, NM_00129778.1:c.710G>T, NM_00129778.1:c.710G^T, NM_00129778.1:c.710G^T, NM_00129778.1	Leber congenital amaurosis type 9 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the NMNATI gene located on chromosomal region 1p36.22. The age of onset is early infantile. This disease is characterized by blindness, nystagmus, roving eye movement,	600,25
NPC1	Niemann-Pick disease, type Cl	NM_000271.4	NM_000271.4:c.33662delT, NM_000271.4:c.3611_3614delTTAC, NM_000271.4:c.3467A>G, NM_000271.4:c.3425T>C, NM_000271.4:c.3425T>C, NM_000271.4:c.3105C>T, NM_000271.4:c.3105C>T, NM_000271.4:c.3105C>T, NM_000271.4:c.3105C>T, NM_000271.4:c.3010C>T, NM_000271.4:c.3010C>T, NM_000271.4:c.3010C>T, NM_000271.4:c.2972_2973delAG, NM_000271.4:c.2932_2973delAG, NM_000271.4:c.2932_2973delAG, NM_000271.4:c.2932_2973delAG, NM_000271.4:c.2932_2973delAG, NM_000271.4:c.2932_2973delAG, NM_000271.4:c.3010C>T, NM_000271.4:c.3	leading to severe visual impairment. Niemann-Pick disease type C1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the NPC1 gene located on chromosomal region 18q11.2. The age of onset varies between the perinatal period and the age of 50 years or more. This disease is characterized by hepatosplenomegaly and progressive neurological involvement. The prevalence is 1/130,000. Niemann-Pick disease	600,25
NPC2	Niemann-pick disease, type C2	NM_006432.3	NM_006432.3:c.436C>T, NM_006432.3:c.358C>T, NM_006432.3:c.358C>T, NM_006432.3:c.355C>T, NM_006432.3:c.355C>T, NM_006432.3:c.355C>A, NM_006432.3:c.35C>A, NM_006432.3:c.27delG	type C2 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the NPC2 gene located on chromosomal region 14q24.3. The age of onset varies between the perinatal period and the age of 50 years or more. This disease is characterized by hepatosplenomegaly and progressive neurological	600,25
NРНР3	Meckel syndrome type 7	NM_153240.4	NM_153240.4:c.3406C>T, NM_153240.4:c.3373C>T, NM_153240.4:c.3576dupa, NM_153240.4:c.2594-1.2_c694-1delAG, NM_153240.4:c.2594-2.2-G94-1delAG, NM_153240.4:c.2591>-C, NM_153240.4:c.25691>-C, NM_153240.4:c.1381G>T, NM_153240.4:c.1381	involvement. The prevalence is 1/130,000. Meckel syndrome type 7 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the NPHP3 gene located on chromosomal region 3q22.1. The age of onset is infantile. This is a disorder	600,25
NPHP4	Nephronophthisis type 4	NM_015102.4	NM_015102.4c:3767_3768insAA, NM_015102.4c:3231-1G>C, NML_015102.4c:2940_2944dupGCTCC, NM_015102.4c:2335C>T, NM_015102.4c:1972C=T, NM_015102.4c:1120-1G>C, NML015102.4c:1120-1G>C, NML015102.4c:1120-1G-C, NML015102.4c:1120-1G	infantile. This disease	600,25

			NM_DIJIOZACIJOCARIJI, NM_DIJIOZACIJI (* 1	is a progressive tubulo- interstitial kidney disorder characterized by polydipsia, polyuria, anemia and growth retardation. The prevalence is 13,000,000. Nephrotic syndrome type 1 follows an autosomal	ı
NPHSI	Nephrotic syndrome, type	I NM_004646.3	NM_004646.3:c.3478C>T, NM_004646.3:c.3325C>T, NM_004646.3:c.350dupG, NM_004646.3:c.3550delG, NM_004646.3:c.359delG, NM_004646.3:c.2928C>T, NM_004646.3:c.2928C>T, NM_004646.3:c.1715C>A, NM_004646.3:c.1481delC, NM_004646.3:c.1307_1308dupAC, NM_004646.3:c.12]_122delCT	recessive pattern of inheritance and is caused by pathogenic variants in the NPHSI gene located on chromosomal region 19q13.12. The age of onset is fetal- infantile. This disease is characterized by fetal proteinuria and nephritic infantile syndrome. The prevalence is 1 in 8 200 births. Enhaced S-Cone	
NR2E3	Enhanced S-cone syndrome	NM_014249.3	NM_014249.3:c.119-2A>C, NM_014249.3:c.226C>T, NM_014249.3:c.298.299deITG, NM_014249.3:c.932G>A, NM_014249.3:c.1034_1038deITGCAG	Syndrome follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the NR2E3 gene located on chromosomal region 15q23. The age of onset is infantile. This disease is characterized by night blindness, reduced bilateral visual acuity, and typical fundus findings (progressive pigmentary degenerative changes, macular edema,	600,25
OCA2	Oculocutaneous albinism type 2	NM_000275.2	NM_000275.2:c.2228C>T, NM_000275.2:c.1960delG, NM_000275.2:c.1842*IG>T, NM_000275.2:c.1465A>C, NM_000275.2:c.1364*IG>T, NM_000275.2:c.1327G>A, NM_000275.2:c.1382*2T>C, NM_000275.2:c.132G>A, NM_000275.2:c.1025A>G, NM_000275.2:c.819_822delCTGGinsGGTC, NM_000275.2:c.157delA, NM_000275.2:c.79G>A	retinoschisis). Oculocutaneous albinism type 2 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the OCA2 gene located on chromosomal region 15q12-q13. The age of onset is infantile. This disease is characterized by variable hypopigmentation of the skin and hair, numerous characteristic ocular changes and misrouting of the optic nerves at the chiasm. The prevalence is	600.25
ОТОА	Deafness, autosomal recessive type 22	NM_144672.3	NM_144672.3:c:121-1G>A, NM_144672.3:c:828deIT, NM_144672.3:c:1725_1726delCA	1/38,000-1/40,000 Deafness, autosomal recessive type 22 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the OTOA gene located on chromosomal region 16p12.2. The age of onset is infantile. This disease is characterized by hearing loss with no associated visible abnormalities of the external ear or any related medical problems.	600,25
OTOF	Auditory neuropathy, autosomal recessive, type 1	NM_001287489.1	NM_001287489.1:c.5474_5475delCC, NM_001287489.1:c.5473C>G, NM_001287489.1:c.5173C>G, NM_001287489.1:c.5103-127>A, NM_001287489.1:c.353C3P>C, NM_001287489.1:c.303Z7>C, NM_001287489.1:c.2348delG, NM_001287489.1:c.178delT, NM_001287489.1:c.15447>C, NM_001287489.1:c.178delT, NM_001287489.1:c.166-2A>G, NM_001287489.1:c.166-2A>G, NM_001287489.1:c.149G>A	Auditory neuropathy, autosomal recessive type 1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the OTOF gene located on chromosomal region 2p23.3. Patients can have varying degrees of hearing loss with poor speech reception out of proportion to the degree	600,25
OTOF	Auditory neuropathy, autosomal recessive, type 1	NM_004802.3	NM_0048023:c.3515G>A	of hearing loss. Auditory neuropathy, autosomal recessive type 1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the OTOF gene located on chromosomal region 2p23.3. Patients can have varying degrees of hearing loss with poor speech reception out of proportion to the degree of hearing loss.	600,25
РАН	Phenylketonuria	NM_000277.1	NM_0002771:c.1315+1G>A, NM_0002771:c.1243G>A, NM_0002771:c.1241A-G, NM_0002771:c.1286-C, NM_0002771:c.1286-C, NM_0002771:c.1286-C, NM_0002771:c.1286-C, NM_0002771:c.1286-C, NM_0002771:c.1286-C, NM_0002771:c.1974-T, NM_0002771:c.1991-17-C, NM_0002771:c.1991-17-C, NM_0002771:c.1991-17-C, NM_0002771:c.1991-17-C, NM_0002771:c.1961-C, NM_0002771:c.1664-C, NM_0002771:c.1664-C, NM_0002771:c.1664-C, NM_0002771:c.1045-TC, NM_0002771:c.1045-TC, NM_0002771:c.1045-TC, NM_0002771:c.1045-TC, NM_0002771:c.1045-TC, NM_0002771:c.1045-TC, NM_0002771:c.1045-C, NM_0002771:c.1050-C, NM_0002771:c.1050-C, NM_0002771:c.1050-C, NM_0002771:c.1050-C, NM_0002771:c.1050-C, NM_0002771:c.1050-C, NM_0002	Phenylketonuria follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PAH gene located on chromosomal region 12q23.2. The age of onset is neonatal. This disease is characterized by gradual developmental delay, stunted growth, microcephaly, seizures, tremors, eczema, vomiting, and musty odor. Untreated patients subsequently develop	600,25

NM_000277.1:c.441+1G>A, NM_000277.1:c.357delC,
NM_000277.1:c.331C>T, NM_000277.1:c.320A>G, NM_000277.1:c.331C>A,
NM_000277.1:c.284_286delTCA, NM_000277.1:c.26LC>A,
NM_000277.1:c.25Go>T, NM_000277.1:c.264A>T, NM_000277.1:c.194T>C,
NM_000277.1:c.165T>G, NM_000277.1:c.143T>C, NM_000277.1:c.136G>A,
NM_000277.1:c.117C>G, NM_000277.1:c.47_48delCT

			NM_000277.I:c.I651>G, NM_000277.I:c.I431>C, NM_000277.I:c.I36G>A, NM_000277.I:c.I37C>G, NM_000277.I:c.47_48delCT	
PANK2	Neurodegeneration with brain iron accumulation type 1	NM_153638.3	NM_153638.3:c.790C>T, NM_153638.3:c.823_824delCT, NM_153638.3:c.1561G>A, NM_153638.3:c.1583C>T	Neurodegeneration with brain iron accumulation type I follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PANK2 gene located on chromosomal region 20pl3. The age of onset is infantile. This disease is characterized by progressive extrapyramidal dysfunction (dystonia, rigidity, choreoathetosis), iron accumulation on the brain and axonal spheroids in the central nervous system. The prevalence is 1-2/1,000,000.
PC	Pyruvate carboxylase deficiency	NM_000920.3	NM_000920.3:c:1748G>T, NM_000920.3:c:434T>C	Pyruvate carboxylase deficiency follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PC gene located on chromosomal region 11q13.2. The age of onset is infantile. This disease is characterized by metabolic acidosis, failure to thrive, developmental delay, and recurrent seizures. The prevalence is 1255,000.
PCCA	Propionic acidemia	NM_000282.3	NM_000282.3:c.229C>T, NM_000282.3:c.261dupT, NM_000282.3:c.600+1G>A, NM_000282.3:c.61023dupT, NM_000282.3:c.1023dupT, NM_000282.3:c.1023dupT, NM_000282.3:c.1023dupT, NM_000282.3:c.1023dupT, NM_000282.3:c.1226_1227deITT, NM_000282.3:c.128416>C, NM_000282.3:c.128416>C, NM_000282.3:c.1899+4_1899+7deIAGTA	Propionic acidemia follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PCCA gene located on chromosomal region 13q32.3. The age of onset is infantile. This disease is characterized by life threatening episodes of metabolic decompensation, neurological dysfunction and may be complicated by cardiomyopathy. The prevalence is 1100,000.
РССВ	Propionic acidemia	NM_001178014.1	NM_001178014.1c.331C>T, NM_001178014.1c.337C>T, NM_001178014.1c.632C>T, NM_001178014.1c.622C>A, NM_001178014.1c.73C>T, NM_001178014.1c.73C>T, NM_001178014.1c.73C>T, NM_001178014.1c.73C>T, NM_001178014.1c.73C>T, NM_001178014.1c.1278.1c.73C>T, NM_001178014.1c.1279.1284de1GCCATCATCCGCGCInsTAGAGCACAGGANM_001178014.1c.1281.2c.73C>T, NM_001178014.1c.1288C>T, NM_001178014.1c.1364C>T, NM_001178014.1c.1364A>C, NM_001178014.1c.1343C>T, NM_001178014.1c.1364A>C, NM_001178014.1c.1364C>T, NM_001178014.1c.1364A>C, NM_001178014.1c.1364C>T, NM_001178014.1c.1364A>C, NM_001178014.1c.1364C>T, NM_001178014.1c.1364A>C	Propionic acidemia follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PCCB gene located on chromosomal region 302.3. The age of onset is infantile. This disease is characterized by life threatening episodes of metabolic decompensation, neurological dysfunction and may be complicated by cardiomyopathy. The prevalence is 1100,000. Deafness, autosomal
PCDH15	Deafness, autosomal recessive type 23	NM_001142763.1	NM_001142763.1c.5680A>T, NM_001142763.1c.4982_4983insTGAT, NM_001142763.1c.4988_4961dupTGAT, NM_001142763.1c.4885delA, NM_001142763.1c.4985.4961dupTGAT, NM_001142763.1c.3733-2A>G, NM_001142763.1c.26661delAT, NM_001142763.1c.1955C>G, NM_001142763.1c.1955C>G, NM_001142763.1c.1955C>G, NM_001142763.1c.10326T, NM_001142763.1c.10326T, NM_001142763.1c.415C>T, NM_001142763.1c.415C_T, NM_00114276	recessive 23 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PCDHIS gene located on chromosomal region loq211. This is a form of non-syndromic 600,25 sensorineural hearing loss. Sensorineural deafness results from damage to the neural receptors of the inner ear, the nerve pathways to the brain, or the area of the brain that receives sound
PDE6A	Retinitis pigmentosa type 43	NM_000440.2	NM_000440.2:c.2053G>A, NM_000440.2:c.1749C>G, NM_000440.2:c.1683G>A, NM_000440.2:c.1560dupA, NM_000440.2:c.1113+1G>T, NM_000440.2:c.1113+1G>A	information. Retinitis pigmentosa type 43 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PDE6A gene located on chromosomal region 532. The age of onset is variable. This disease is characterized by night blindness (nyctalopia) due 600,25 to loss of rod function, often in adolescence or earlier. They then develop peripheral visual field impairment, and overtime loss of central vision, usually at late stages, often around midlife. The prevalence is 1:10,000-
PDE6B	Retinitis pigmentosa type 40	NM_000283.3	NM_000283.3'c.892C>T, NM_000283.3'c.1540delC, NM_000283.3'c.1572delC, NM_000283.3'c.1580T>C, NM_000283.3'c.1669C>T, NM_000283.3'c.1920+2T>C	\$10,000. Retinitis pigmentosa 40 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PDE68 gene located on chromosomal region 4p16.3. The age of onset is variable. Retinitis pigmentosa 40 is a retinal 600,25 dystrophy belonging to the group of pigmentary retinopathies. This disease is characterized by night bilindness, followed by a progressive loss of peripheral vision in the

Р	PEXI	Heimler syndrome type 1	NM_000466.2	NM_000466.2:c.3505_3517delCAGTTGTTTCAC, NM_000466.2:c.2516delA, NM_000466.2:c.2528G>A, NM_000466.2:c.2057dupt; NM_000466.2:c.1991T>C, NM_000466.2:c.1952_1960dupCAGTGTGGA, NM_000466.2:c.1842delA, NM_000466.2:c.1239+1G>T, NM_000466.2:c.877C>T	daylight period and leading to blindness. Heimler syndrome I follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PEXI gene located on chromosomal region 7q212. This disease is characterized by sensorineural hearing loss, enamel hyoplasia of the secondary dentition, and nail abnormalities. Rhizomelic chondrodysplasia is punctata type I follows an	600,25
P	PEX7	Rhizomelic chondrodysplasia punctata, type 1	NM_000288.3	NM_000288.3c.532C>T, NM_000288.3c.618G>A, NM_000288.3c.649G>A, NM_000288.3c.653C>T, NM_000288.3c.694C>T, NM_000288.3c.854A>G, NM_000288.3c.875T>A, NM_000288.3c.903+1G>C	autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PEX7 gene located on chromosomal region 6q253. The age of onset is early. This disease is characterized by proximal shortening of the humerus and to a lesser degree the femur (rhizomelia), punctate calcifications in cartilage with epiphyseal and metaphyseal abnormalities (chondrodysplasia punctata), coronal clefts of the vertebral bodies, cataracts, postnatal growth deficiency is profound, intellectual disability is severe, seizures. The prevalence is	600,25
Р	РНΥН	Refsum disease	NM_001323082.1	NM_001323082.hc.830G>A, NM_001323082.hc.829C>T, NM_001323082.hc.829C>T, NM_001323082.hc.829C>T, NM_001323082.hc.684+5C>T, NM_001323082.hc.503-2A>G, NM_001323082.hc.135-1G>C, NM_001323082.hc.135-1G>C, NM_001323082.hc.135-1G>C, NM_001323082.hc.135-2A>G	<1:100,000 Refsum disease follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PHYH gene located on chromosomal region 10pl3. The age of onset is variable. This disease is characterized by hemeralopia (loss of vision in the dark), followed by episods of chronic distal motor polyneuropathy. Other associated signs include perceptive deafness, anosmia, cerebellous ataxia and sometimes, severe intellectual deficiency. Over the course of time cutaneous signs appear (ichtyosis), along with polyepiphyseal dysplasia, myocardiopathy, elevated protein in cerebrospinal fluid, and pigmentary retinitis that may result in blindness. The prevalence is 11.000,000-91.000,000.	600,25
P	PKHDI	Polycystic kidney disease type 4	NM_138694.3	NM_138694_3:c.12027C>G, NM_138694_3:c.1363_11372delCTTCCCTGGA, NM_138694_3:c.10452dupT, NM_138694_3:c.10412T>G, NM_138694_3:c.10452dupT, NM_138694_3:c.97913C>A, NM_138694_3:c.953017>C, NM_138694_3:c.953017>C, NM_138694_3:c.953017>C, NM_138694_3:c.953017>C, NM_138694_3:c.84076>C, NM_138694_3:c.8802d>C, NM_138694_3:c.88070>C, NM_138694_3:c.8802d>C, NM_138694_3:c.88070>T, NM_138694_3:c.53765_325_5326_61AG, NM_138694_3:c.3766_61C, NM_138694_3:c.3595_5326_61AG, NM_138694_3:c.3576_54C, NM_138694_3:c.3525_5326_61AG, NM_138694_3:c.3576_54C, NM_138694_3:c.35229>2AP NM_38694_3:c.3529>2AP NM_38694_3:c.3576_52C, NM_38694_3:c.3576_5A, NM_38694_3:c.3259>2AP NM_38694_3:c.2829>2AP NM_38694_3:c.2452>T, NM_38694_3:c.2452>T, NM_38694_3:c.2462>T, NM_38694_3:c.2462>T, NM_38694_3:c.3576_52 NM_38694_3:c.1462>T, NM_38694_3:c.3462>T, NM_38694_3:c.3562>AP NM_38694_3:c.1462>T, NM_38694_3:c.3546>T, NM_38694_3:c.3562>AP NM_38694_3:c.1462>T, NM_38694_3:c.370<>T, NM_38694_3:c.35046C, NM_138694_3:c.370<>T, NM_38694_3:c.35046C, NM_138694_3:c.370<>T, NM_138694_3:c.35046C, NM_138694_3:c.370<>T, NM_138694_3:c.35046C, NM_138694_3:c.370<>T, NM_138694_3:c.35046C, NM_138694_3:c.370< <t, nm_138694_3:c.35046c,="" nm_138694_3:c.370<<t,="" nm_138694_3:c.3<="" td=""><td>autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PKHD1 gene located on chromosomal region 6p12.3-p12.2 The age of onset is early. This disease is a severe form of polycystic kidney disease affecting the kidneys and, in some cases, the hepatic biliary tract. Up to 50% of the affected neonates die shortly after birth, as a result of severe pulmonary hypoplasia and secondary respiratory insufficiency. In the subset that survives the perinatal period, morbidity and mortality are mainly related to severe systemic hypertension, renal insufficiency, and portal hypertension due to</td><td>600,25</td></t,>	autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PKHD1 gene located on chromosomal region 6p12.3-p12.2 The age of onset is early. This disease is a severe form of polycystic kidney disease affecting the kidneys and, in some cases, the hepatic biliary tract. Up to 50% of the affected neonates die shortly after birth, as a result of severe pulmonary hypoplasia and secondary respiratory insufficiency. In the subset that survives the perinatal period, morbidity and mortality are mainly related to severe systemic hypertension, renal insufficiency, and portal hypertension due to	600,25
Р	KLR	Pyruvate kinase deficiency	NM_000298.5	NM_000298.5:c.1675C>T, NM_000298.5:c.1529G>A, NM_000298.5:c.1528C>T, NM_000298.5:c.1456C>T, NM_000298.5:c.1251C>A, NM_000298.5:c.1251C>A, NM_000298.5:c.1151C>T, NM_000298.5:c.721G>T	portal-tract fibrosis. Pyruvate kinase deficiency follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PKLR gnee located on chromosomal region 1q22. The age of onset is early. This disease is characterized by highly variable degree of chronic hemolysis, with severe neonatal jaundice and fatal anemia at birth, severe transfusion-dependent chronic hemolysis, and moderate hemolysis with exacerbation during infection. The prevalence is 1:20,000. Nephrotic syndrome type 3 follows an autosomal recessive pattern of inheritance and is caused.	600,25
Р		Nephrotic syndrome, type 3	NM_016341.3	NM_016341.3:c.961C>T, NM_016341.3:c.3346C>T, NM_016341.3:c.3736C>T, NM_016341.3:c.3846delC, NM_016341.3:c.4451C>T, NM_016341.3:c.4809delA, NM_016341.3:c.5560C>T	inheritance and is caused by pathogenic variants in the PLCEI gene located on chromosomal region 10q23.33. The age of onset is variable. This disease is characterized by low blood	

PLG	Plasminogen deficiency, type I	NM_000301.3	NM_000301.3:c:112A>G, NM_0003013:c:693_695delGAA, NM_000301.3:c:704G>A, NM_0003013:c:1120G>T, NM_000301.3:c:1435G>T, NM_000301.3:c:1848G>A	protein levels, high cholesterol levels, high cholesterol levels, and presence of protein in the urine. The prevalence is 2:100,000-7:100,000 Children; 3:100,000 adults. Plasminogen deficiency type I follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PLG gene located on chromosomal region 6q.26. The age of onset is infantile. This disease is characterized by markedly impaired extracellular fibrinolysis leading to the formation of ligneous (fibrin-rich) pseudomembranes on mucosae during wound healing. The prevalence is 11,000,000-91,000,000. Congenital disorder of glycosylation type Ia follows an autosomal recessive pattern of	600,25
РММ2	Congenital disorder of glycosylation, type la	NM_000303.2	NM_000303.2c.26G>A, NM_000303.2c.53C>G, NM_000303.2c.95T>G, NM_000303.2c.995_96delTAinsGC, NM_000303.2c.97C>T, NM_000303.2c.195C>T, NM_000303.2c.13T>C, NM_000303.2c.199C>T, NM_000303.2c.13T>C, NM_000303.2c.190delT NM_000303.2c.35G>T, NM_000303.2c.35T>C, NM_000303.2c.35G>T, NM_000303.2c.36G9T>C, NM_000303.2c.3C.3C.3C.3C.3C.3C.3C.3C.3C.3C.3C.3C.3C.	inheritance and is caused by pathogenic variants in the PMM2 gene located on chromosomal region [6p13.2. The age of onset is infantile. This disease is characterized by highly variable clinical manifestations that may include feeding problems, womiting, and diarrhea with failure to thrive in infants, and severe encephalopathy with axial hypotonia, abnormal eye movement, marked psychomotor retardation, peripheral neuropathy, cerebellar hypoplasia, stroke-like episodes, and retinitis pigmentosa in late infancy, childhood or adulthood.	600,25
POLG	Mitochondrial DNA depletion syndrome 4A (Alpers type)	NM_001126131.1	NM_001126131.1c.3644-1G>A, NM_001126131.1c.3630dupC, NM_001126131.1c.3286C>T, NM_001126131.1c.3218C>T, NM_001126131.1c.2794C>T, NM_001126131.1c.2794C>T, NM_001126131.1c.2605C>T, NM_001126131.1c.2591A>G, NM_001126131.1c.2565C>T, NM_001126131.1c.2591A>G, NM_001126131.1c.254G>C, NM_001126131.1c.254G>A, NM_001126131.1c.254GAA, NM_001126131.1c.254GAA, NM_001126131.1c.254GAA, NM_001126131.1c.254GAA, NM_001126131.1c.254GAA, NM_001126131.1c.254GAA, NM_001126131.1c.254GAA, NM_001126131.1c.2	Mitochondrial DNA depletion syndrome, Alpers type follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the POLG gene	600,25
POMGNT	Muscular dystrophy- q dystroglycanopathy (congenital with brain and eye anomalies), type A, 3	NM_001243766.1	NM_001243766.1:c.1864delC, NM_001243766.1:c.1814G>C, NM_001243766.1:c.1864delC, NM_001243766.1:c.1539+IG>T, NM_001243766.1:c.1539+IG>T, NM_001243766.1:c.1549G-A, NM_001243766.1:c.141A>T, NM_001243766.1:c.141A>T, NM_001243766.1:c.141A>T, NM_001243766.1:c.141A>T, NM_001243766.1:c.193Co-A, NM_001243766.1:c.193Co-A, NM_001243766.1:c.593Co-A, NM_001243766.1:c.593Co-T, NM_001243766.1:c.593Co-T, NM_001243766.1:c.92dupA	(congenital with brain and eye anomalies) type A3 which includes both the more severe Walker-Warburg syndrome (WWS) and the slightly less severe muscle-eyebrain disease (MEB), follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the POMGNTI gene located on chromosomal region Ip34-1. The age of onset is infantile. This disease is characterized by generalized severe hypotonia, muscle weakness, absent psychomotor development, eye involvement and seizures.	600,25
РОМТІ	Muscular dystrophy- dystroglycanopathy (congenital with brain and eye anomalies), type A, 1	NM_007171.3	NM_007171.3:c.193G>A, NM_007171.3:c.226G>A, NM_007171.3:c.598G>C, NM_007171.3:c.193G>T, NM_007171.3:c.831C>G, NM_007171.3:c.907C>T, NM_007171.3:c.183G>T, NM_007171.3:c.1242-2A>G, NM_007171.3:c.1261dpc, NM_007171.3:c.1280_1281delAGinsTC, NM_007171.3:c.1280_1281delAGinsTC, NM_007171.3:c.1545C>G, NM_007171.3:c.146G>C, NM_007171.3:c.2163C>A, NM_007171.3:c.2163C>A, NM_007171.3:c.2167dupG	The prevalence is 1-9100,000. Muscular dystrophy-dystroglycanopathy (congenital with brain and eye anomalies), type Al follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the POMTI gene located on chromosomal region 9354.13. Congenital muscular dystrophy-dystroglycanopathy with brain and eye anomalies (type Al, which includes both the more severe Walker-Warburg syndrome (WWS) and the slightly less severe muscle-eye-brain disease (MEB), is a genetically heterogeneous disorder with characteristic brain and eye malformations, profound mental retardation, congenital muscular dystrophy, and early death. The phenotype commonly includes cobblestone (type II) lissencephaly, cerebellar malformations. Muscular dystrophy-dystroglycanopathy (congenital with brain and	600,25

РОМТ2	Muscular dystrophy- dystroglycanopathy (congenital with brain and eye anomalies), type A, 2	NM_013382.5	NM_013382.5:c.2243G>C, NM_013382.5:c.2177G>A, NM_013382.5:c.1994G>A, NM_013382.5:c.1994G>A, NM_013382.5:c.1926-2A>G, NM_013382.5:c.1608_1609delCA, NM_013382.5:c.1645G>T, NM_013382.5:c.1608_1609delCA, NM_013382.5:c.1445G>T, NM_013382.5:c.1045_056666666666666666666666666666666666	follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the POMT2 gene located on chromosomal region 14q24-3. Congenital muscular dystrophydystroglycanopathy with brain and eye anomalies (type A), which includes both the more severe Walker-Warburg syndrome (WWS) and the slightly less severe muscle-eye-brain disease (MEB), is a genetically heterogeneous disorder with characteristic brain and eye malformations, profound mental retardation, congenital muscular dystrophy, and early death. The phenotype commonly includes cobblestone (type II) lissencephaly, cerebellar malformations, and retinal malformations.	600,25
РРП	Ceroid lipofuscinosis, neuronal, type 1	NM_000310.3	NM_0003103:c.840dupA, NM_000310.3:c.627+1G>T, NM_000310.3:c.54IG>T, NM_000310.3:c.45IC>T, NM_000310.3:c.223A>C, NM_000310.3:c.169dupA, NM_000310.3:c.29T>A	lipofuscinoses, type 1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the PPTI gene located on chromosomal region 1p32.	600,25
PROMI	Retinitis pigmentosa, type 41	NM_006017.2	NM_006017.2:c.2490-2A>G, NM_006017.2:c.1841delG, NM_006017.2:c.1726C>T, NM_006017.2:c.1354dupT, NM_006017.2:c.1177_1178delAT, NM_006017.2:c.199C>T	inheritance and is caused by pathogenic variants in the PROMI gene located on chromosomal region 4p15.32. The age of onset is early. This disease is characterized by night blindness often in adolescence or earlier. They then develop peripheral visual field impairment, and overtime loss of central vision, usually at late stages, often around midlife. The prevalence is 19:100,000-	600,25
PYGM	McArdle disease	NM_005609.3	NM_005609.3:c.2392T>C, NM_005609.3:c.2262delA, NM_005609.3:c.128.2130delTTC, NM_005609.3:c.1963G>A, NM_005609.3:c.1282-1350delTTC, NM_005609.3:c.1768-16-A, NM_005609.3:c.1768-16-A, NM_005609.3:c.1762T-C, NM_005609.3:c.1623G>T, NM_005609.3:c.1623G>T, NM_005609.3:c.1623G>T, NM_005609.3:c.1523G>T, NM_005609.3:c.255C>A, NM_005609.3:c.255C>A, NM_005609.3:c.148C>T, NM_005609.3:c.13_14delCT, NM_005609.3:c.13_14delCT, NM_005609.3:c.13_14delCT, NM_005609.3:c.15	onset is infantile. This disease is characterized by muscular exercise intolerance with myalgia, cramps, fatigue, and muscle weakness.	
RAGI	Omenn syndrome; Severe combined immunodeficiency, B cell- negative	NM_000448.2	NM_000448.2:c.256_257delAA, NM_000448.2:c.940C>T, NM_000448.2:c.983G>A, NM_000448.2:c.168IC>T, NM_000448.2:c.1682G>A, NM_000448.2:c.2164G>A, NM_000448.2:c.235G>T, NM_000448.2:c.2814T>G, NM_000448.2:c.2923C>T	Omenn syndrome follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the RAG1 and RAG2 genes located on chromosomal region Ilp12. The age of onset is early. This disease is characterized by erythroderma, desquamation, alopecia, chronic diarrhea, failure to thrive, lymphadenopathy, and hepatosplenomegaly, associated with severe combined immunodeficiency, Severe combined immunodeficiency, autosomal recessive, Tell-negative (F.), B cell negative (B-), NK cell positive (IKH+) is also caused by mutation in the RAG1 and RAG2 genes. This disease is characterized by impairment of both humoral and cell-mediated immunity, leukopenia, and low or absent antibody levels. Patients present in infancy recurrent, persistent infections by opportunistic organisms. The common characteristic of all types of SCID is absence of T-cell-mediated cellular immunity due to a defect in T-cell development. Without treatment, patients usually die within the first year of life. Fetal akinesia deformation sequence follows an	600,25
			NM 005055.4:c.848T>C. NM 005055.4:c.807C>A.	autosomal recessive pattern of inheritance and is caused by pathogenic variants in the RAPSN	

RAPSN	Fetal akinesia deformation sequence	NM_005055.4	NM_005055.4:c:566C>T, NM_005055.4:c:490C>T, NM_005055.4:c:484G>A, NM_005055.4:c:416T>C, NM_005055.4:c:264C>A	gene located on chromosomal region 600,25 Ilp11.2. The age of onset is early. This disease is characterized by multiple joint contractures, facial anomalies and pulmonary hypoplasia. The
RAX	Isolated microphthalmia, type 3	NM_013435.2	NM_013435.2:c.909C>G, NM_013435.2:c.439C>T, NM_013435.2:c.383_384delAG, NM_013435.2:c.18C>A	prevalence is 13,000. Isolated microphthalmia type 3 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the RAX gene located on chromosomal region 18q21.32. Microphthalmia designates a heterogeneous group of coular malformations with a more or less evident reduction in the size of the eyeball. Additional features include high hypermetropia and a short axial length.
RDH12	Leber congenital amaurosis, type 13	NM_152443.2	NM_152443.2:c.146C>T, NM_152443.2:c.152T>A, NM_152443.2:c.184C>T, NM_152443.2:c.210dupC, NM_152443.2:c.295C>A, NM_152443.2:c.379C>T, NM_152443.2:c.379C-T,	characterized by blindness, nystagmus, roving eye movement and lack of detectable signals on an electroretinogram, leading to severe visual impairment within the first year of life.
RGR	Retinitis pigmentosa, type 44	NM_002921.3	NM_002921.3:c.262_269dupGGCTCGGA, NM_002921.3:c.273_274insGGCTCGGA, NM_002921.3:c.877C>T	Retinitis pigmentosa type 44 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the RGR gene located on chromosomal region 10q23.1. The age of onset is variable. This disease is characterized by night blindness (nyctalopia) due 600,25 to loss of rod function, often in adolescence or earlier. They then develop peripheral visual field impairment, and overtime loss of central vision, usually at late stages, often around midlife. The prevalence is 110,000- \$10,000.
RHO	Retinitis pigmentosa, type 4, autosomal recessive	NM_000539.3	NM_000539.3:c:173C>T, NM_000539.3:c:448G>A, NM_000539.3:c:620T>G, NM_000539.3:c:745G>T	Retinitis pigmentosa type 4 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the RHO gene located on chromosomal region 3q221. The age of onset is variable. This disease is characterized by night blindness (nyctalopia) due 600,25 to loss of rod function, often in adolescence or earlier. They then develop peripheral visual field impairment, and overtime loss of central vision, usually at late stages, often around midlife. The prevalence is 1:10,000-5:10,000.
RLBPI	Bothnia retinal dystrophy	NM_000326.4	NM_000326.4:c.700C>T; NM_000326.4:c.452G>A, NM_000326.4:c.333T>G	Bothnia retinal dystrophy follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the RLBP1 gene located on chromosomal region 15q26.1. This disease is characterized by night blindness from early childhood with features consistent with retinitis punctata albescens and macular degeneration. The prevalence is unknown.
RPE65	Leber congenital amaurosis, type 2	NM_000329.2	NM_000329.2:c.1543C>T, NM_000329.2:c.1355T>G, NM_000329.2:c.1292A>G, NM_000329.2:c.1102T>C, NM_000329.2:c.10287C>A, NM_000329.2:c.1067delA, NM_000329.2:c.10287C>C, NM_000329.2:c.907A>T, NM_000329.2:c.514_515delGT, NM_000329.2:c.271C>T	Leber congenital amaurosis 2 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the RPEGS gene located on chromosomal region 1p31.3-p31.2. The age of onset is variable. This disease is characterized by a severe dystrophy of the retina, typically becoming evident in the first years of life. Visual function is usually poor and often accompanied by nystagmus, sluggish or near-absent pupillary responses, photophobia, high hyperopia and keratoconus. Joubert syndrome (JBTS) type 7 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the RPGRIPIL canal located on

chromosomal region 16q12.2. The age of onset is early. JBTS is characterized by congenital malformation of the brainstem and agenesis of the cerebellar vermis

(molar tooth sign) leading to an abnormal respiratory pattern, nystagmus, hypotonia, mental retardation, ataxia, and delay in achieving motor milestones. Other variable features include retinal

features include retinal dystrophy (less common in JBTS7) and nephronophthisis (usually juvenile). The prevalence is 1:100,000. RPGRIPIL gene

600.25

is also associated with Meckel syndrome type 5, a rare, autosomal recessive lethal condition characterized by central

nervous system polydactyly, multicystic kidney dysplasia, and kidney dysplasia, and ductal proliferation in the portal area of the liver. Other phenotype associated is COACH syndrome, an autosomal

areas of reduced mitochondrial oxidative

show sarcomere

disorganization and

including neonatal

progress slowly or remain

600.25

ophthalmoplegia, although there is

although there is phenotypic variability. Some patients may present in utero with fetal akinesia, arthrogryposis, and lung hypoplasia resulting in fetal or perinatal death (McKie et al., 2014). Skeletal muscle historic architecture of a patients with the surface of a patients with the surface of a patients with a perinatal death (McKie et al., 2014). Skeletal muscle

(Jungbluth et al., 2007), congenital fiber-type disproportion (CFTD) (Monnier et al., 2009), and centronuclear myopathy (Wilmshurst et al., 2010).

Spastic ataxia. Charlevoix Saguenay type follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SACS gene located on chromosomal 600.25

The prevalence is 1:1,500-

malformations, postaxial,

recessive disorder characterized by mental retardation, ataxia due to cerebellar hypoplasia, and hepatic fibrosis. Other features, such as coloboma and renal cysts,

coloboma and renal cysts, may be variable. COACH syndrome is considered by some to be a subtype of Joubert syndrome with congenital hepatic fibrosis.

Multiminicore disease (MMD) is an inherited neuromuscular disorder defined pathologically by the presence of multiple

mitochondrial oxidative activity running along a limited extent of the longitudinal axis of the muscle fiber, so-called 'minicores.' These regions

disorganization and mitochondria depletion. Typically, no dystrophic signs, such as muscle fiber necrosis or regeneration or significant endomysial

fibrosis, are present. MMD

is a pathologic diagnosis and shows clinical and genetic heterogeneity. Affected individuals have clinical features of a congenital myopathy,

including neonatal hypotonia, delayed motor development, and generalized muscle weakness and amyotrophy, which may progress slowly or remain

stable (Ferreiro and

stable (Ferreiro and Fardeau, 2002).Patients with recessive mutations in the RYRI gene typically show severe congenital muscular dystrophy with

al, 2014). Skeletal muscle biopsy of patients with recessive RYRI mutations show variable features, including central cores (Jungbluth et al., 2007),

located on chromosoma region 13q11. The age of onset is early. This diseas is characterized by early-onset cerebellar ataxia with spasticity, a pyramidal syndrome and peripheral neuropathy

1:2,000. Oguchi disease type 1 Oguchi disease type I follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SAG gene located on chromosomal region 2g37 The age of onset is infantile. This disease is characterized by congenital stationary

Joubert syndrome, type 7; RPGRIP1L Meckel syndrome, type 5; NM_015272.4 COACH syndrome

NM_015272.4:c.3634_3637delGAAA, NM_015272.4:c.2794_2795delTT, NM_015272.4:c.2614C>T, NM_015272.4:c.2413C>T, NM_015272.4:c.1975T>C, NM_015272.4:c.1975T>C, NM_015272.4:c.1843A>C, NM_015272.4:c.1329dupA, NM_015272.4:c.1326_1329delAAAA, NM_015272.4:c.776+1G>A, NM_015272.4:c.757C>T. NM_015272.4:c.697A>T. NM_015272.4:c.394A>T

DVD1

Minicore myopathy with external ophthalmoplegia NM_000540.2

NM_000540.2:c.325C>T, NM_000540.2:c.487C>T,
NM_000540.2:c.5314ZT>C, NM_000540.2:c.738T>G,
NM_000540.2:c.1021G>A, NM_000540.2:c.138GS¬T,
NM_000540.2:c.1205T>C, NM_000540.2:c.1759_1742dupATCA,
NM_000540.2:c.184G>T, NM_000540.2:c.4076delG,
NM_000540.2:c.449GS-T, NM_000540.2:c.5333C>A,
NM_000540.2:c.45726_5727delAG, NM_000540.2:c.6082C>T, NM_000540_2:c.5726_5727/delAd, NM_000540_2:c.6082C.
NM_000540_2:c.6104A>T. NM_000540_2:c.6721C>T,
NM_000540_2:c.7366C>T, NM_000540_2:c.7300G>A,
NM_000540_2:c.7360C>T, NM_000540_2:c.7337G>A,
NM_000540_2:c.77465_7475delCAAACATCTCAGC,
NM_000540_2:c.7781C>A, NM_000540_2:c.7836-1C>A,
NM_000540_2:c.9000+1G-T, NM_000540_2:c.905dupC,
NM_000540_2:c.9000+1G-T, NM_000540_2:c.905dupC,
NM_000540_2:c.10343C>T, NM_000540_2:c.10579C>T, NM_000540.2:c.13480G>T, NM_000540.2:c.14126C>T, NM_000540.2:c.14365-2A>T, NM_000540.2:c.14545G>A

Spastic ataxia, Charlevoix- NM_014363.5

NM_014363.5:c.13237C>T, NM_014363.5:c.12160C>T NM_0143635:c.8844delT, NM_0143635:c.7504C>T, NM_0143635:c.8564T, NM_0143635:c.6555TA, NM_0143635:c.6555TA, NM_0143635:c.6555C>T, NM_0143635:c.5618_561964LT, NM_0143635:c.4353C>T, NM_0143635:c.3198T>A, NM_0143635:c.994A>T, NM_0143635:c.517C>T

SACS

SAG	Oguchi disease, type 1	NM_000541.4	NM_000541.4:c.298dupG, NM_000541.4:c.523C>T, NM_000541.4:c.577C>T, NM_000541.4:c.874C>-T, NM_000541.4:c.916G>T NM_000541.4:c.926delA, NM_000541.4:c.993C>G	night blindness and the Mizuo-Nakamura phenomenon which is a unique morphological and functional abnormality of the retina that presents with a typical golden- yellow or silver-gray discoloration of the fundus in the presence of light that disappears after dark- adaptation and appears again after the onset of
SBDS	Shwachman-Diamond syndrome	NM_016038.2	NM_016038.2:c.377G>C, NM_016038.2:c.258+2T>C, NM_016038.2:c.184A>T, NM_016038.2:c.183_184delTAinsCT, NM_016038.2:c.120delG	light. Shwachman-Diamond syndrome follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SBDS gene located on chromosomal region 7q11,21. The age of onset is infantile. This disease is characterized by chronic and usually mild neutropenia, pancreatic exocrine insufficiency associated with steatorrhea and growth failure, skeletal dysplasia with short stature, and an increased risk of bone marrow aplasia or leukemic transformation, cutaneous (eczema or ichthyosis) and dental anomalies, and psychomotor retardation. The prevalence is 1:76,000 newborn. Pseudohypoaldosteronism type 1, follows an
SCNNIB	Pseudohypoaldosteronism type 1	^{1,} NM_000336.2	NM_000336.2:c:109G>A	autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SCNNIA (12p13), SCNNIB (16p12, 2p12.1) and SCNNIG (16p12) genes. The age of onset is early. This disease is characterized by severe dehydration, vomiting and failure to thrive occurring in the first weeks of life, the clinical picture may be complicated by cardiac dysrhythmias, collapse,
SCNNIG	Pseudohypoaldosteronism type 1	^{),} NM_001039.3	NM_001039.3:c.600dupA, NM_001039.3:c.1373+2T>C, NM_001039.3:c.1570-1G>A, NM_001039.3:c.1627delG	shock or cardiac arrest. Pseudohypoaldosteronism type 1, follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SCNNIA (12pi3), SCNNIB (16pi2) genes. The age of onset is early. This disease is characterized by severe dehydration, vomiting and failure to thrive occurring in the first weeks of life, the clinical picture may be complicated by cardiac dysrhythmias, collapse, shock or cardiac arrest. Spinocerebellar ataxia
SETX	Spinocerebellar ataxia, autosomal recessive, type	NM_015046.5	NM_015046.5:c.6848_6851delCAGA, NM_015046.5:c.5824_6839delAACAAA, NM_015046.5:c.5927T>G, NM_015046.5:c.5308_6311delGAGA, NM_015046.5:c.4087C>T, NM_015046.5:c.3508_5311delGAGA, NM_015046.5:c.4087C>T, NM_015046.5:c.3022C>T, NM_015046.5:c.166T>C, NM_015046.5:c.1027G>T, NM_015046.5:c.994C>T	with axonal neuropathy type I follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SETX gene located on chromosomal region 9g.3-4.13. The age of onset is infantile. This disease is characterized by 600.25 progressive cerebellar ataxia, axonal sensorimotor neuropathy with oculomotor apraxia, fixation instability, extrapyramidal features and an elevated serum alpha-fetoprotein level. The prevalence is 4:100.000-8:100.000.
SGCA	Muscular dystrophy, limb- girdle, type 2D	NM_000023.3	NM_000023.3:c:101G>A, NM_000023.3:c:229C>T, NM_000023.3:c:371T>C, NM_000023.3:c:518T>C, NM_000023.3:c:573G>A, NM_000023.3:c:850C>T, NM_000023.3:c:903_904dupCC	Autosomal recessive limb- girdle muscular dystrophy type 2D follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SCCA gene located on chromosomal region 4q12. The age of onset is variable. This disease is characterized by limb-girdle weakness and calf pseudohypertrophy. The prevalence is 11,000,000-91,000,000. Autosomal recessive limb-
sgcg	Muscular dystrophy, limb- girdle, type 2C	NM_000231.2	NM_000231.2:c.89delG, NM_000231.2:c.195+4_195+7delAGTA, NM_000231.2:c.505+1G>A, NM_000231.2:c.525delT, NM_000231.2:c.787G>A, NM_000231.2:c.848G>A	girdle muscular dystrophy type 2C follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SGCG gene located on chromosomal region 13q12.12. The age of onset is variable. This disease is characterized by limb-girdle weakness, calf hypertrophy, diaphragmatic weakness, and variable cardiac abnormalities. Mucopolysaccharidosit type 3A (Sanfilippo syndrome type A) follows

SGSH	Mucopolysaccharidosis, type 3A (Sanfilippo A)	NM_000199.3	NM_000199.3:c.1380delT, NM_000199.3:c.1339G>A, NM_000199.3:c.1389G>A, NM_000199.3:c.1376C>A, NM_000199.3:c.1376C>A, NM_000199.3:c.1376C>T, NM_000199.3:c.3576C>T, NM_000199.3:c.456A>T, NM_000199.3:c.456A>T, NM_000199.3:c.456A>A, NM_000199.3:c.356A>A, NM_000199.3:c.356A>A, NM_000199.3:c.356A>A, NM_000199.3:c.350AP, NM_000199	an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SCSH gene located on chromosomal region 17q25.3. The age of onset is infantile. This disease is characterized by behavioural disorders (hyperkinesia, aggressiveness) and intellectual deterioration, sleep disorders and very mild dysmorphism. The prevalence is >1:70,000 newborn.	600,25
SH3TC2	Charcot-Marie-Tooth disease, type 4C	NM_024577.3	NM_024577.3:c.3676-IG-A, NM_024577.3:c.3601C>T, NM_024577.3:c.331delC, NM_024577.3:c.3326C>C, NM_024577.3:c.3326C>C, NM_024577.3:c.2325C>T, NM_024577.3:c.2993_2994insC, NM_024577.3:c.2895_2994insC, NM_024577.3:c.2700-Z, NM_024577.3:c.2491_2492delAG, NM_024577.3:c.2700-Z, NM_024577.3:c.2700-Z, NM_024577.3:c.1982T>C, NM_024577.3:c.1972C>T, NM_024577.3:c.1960-A, NM_024577.3:c.1972C>T, NM_02	Charcot-Marie-Tooth disease, type 4C follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SH3TC2 gene located on chromosomal region 5q32. The age of onset is infantile. This disease is characterized by scollosis or kyphoscollosis, neuropathy, foot deformities, respiratory insufficiency, hypoacousis and deafness. Bartter syndrome type 1 follows an autosomal	
SLC12A1	Bartter syndrome, type 1	NM_000338.2	NM_000338.2c.223C>T, NM_000338.2c.628+2T>C, NM_000338.2c.814G>T, NM_000338.2c.1875G>A, NM_000338.2c.1942G>A, NM_000338.2c.2805dupA, NM_000338.2c.2952_2955delCAAA	recessive pattern of inheritance and is caused by pathogenic variants in the SLC12A1 gene located on chrosomal region	600,25
SLC17A5	Salla disease	NM_012434.4	NM_012434.4:c.1259+1G>A, NM_012434.4:c.406A>G, NM_012434.4:c.115C>T, NM_012434.4:c.43G>T	Salla disease follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SLC17A5 gene located on chromosomal region 6q13.	600,25
SLC26A2	Achondrogenesis, type 1B (diastrophic dysplasia)	NM_000112.3	NM_000112.3:c.496G>A, NM_000112.3:c.532C>T, NM_000112.3:c.833delC, NM_000112.3:c.835c=T, NM_000112.3:c.1020_1022delTGT, NM_000112.3:c.1373A>G, NM_000112.3:c.136A>C, NM_000112.3:c.1373A>G, NM_000112.3:c.1724AelA, NM_000112.3:c.1878delG, NM_000112.3:c.1957T>A, NM_000112.3:c.2033G>T	Actionaryanesis type in (diastrophic dysplasia) follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SLC26A2 gene located on chromosomal region 5.432. The age of onset is early. This disease is characterized by severe micromelia with very short fingers and toes, a flat face, a short neck, thickened soft tissue around the neck, pypollasia of the thorax, protuberant abdomen, a hydropic fetal appearance and distinctive histological features of the cartilage.	
SLC26A4	Deafness, autosomal recessive, type 4	NM_000441.1	NM_000441.tc.269C>T, NM_000441.tc.281C>T, NM_000441.tc.412G>T, NM_000441.tc.254C>C, NM_000441.tc.554T>C, NM_000441.tc.626G>T, NM_000441.tc.626G>T, NM_000441.tc.626G>T, NM_000441.tc.91640µC, NM_000441.tc.9161A>T, NM_000441.tc.1001C>T, NM_000441.tc.1001C>T, NM_000441.tc.103T>C, NM_000441.tc.103T>C, NM_000441.tc.103T>C, NM_000441.tc.103T>C, NM_000441.tc.1054S>C, NM_000441.tc.1054S>C, NM_000441.tc.1054S>C, NM_000441.tc.1256S>C, NM_000441.tc.126S>C, NM_000441.tc.2048T>C, NM_000441.tc.206S>C, NM_000441.tc.206S>C, NM_000441.tc.206S>C, NM_000441.tc.206S>C, NM_000441.tc.206S>C, NM_000441.tc.206S>C, NM_000441.tc.216SC>T, NM_000441.tc.216SA>C	The prevalence is 1:20,000. Autosomal recessive nonsyndromic sensorineural deafness	600,25
SLC37A4	Glycogen storage disease, type lb	NM_001164278.1	NM_001164278.1:c.1309C>T, NM_001164278.1:c.1190-2_1190-1delAG, NM_001164278.1:c.1196_2T, NM_001164278.1:c.1108_G>T, NM_001164278.1:c.1082G>A, NM_001164278.1:c.1081G>T, NM_001164278.1:c.708delGTG, NM_001164278.1:c.552T>C, NM_001164278.1:c.287G>A, NM_001164278.1:c.83G>A	autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SLC37A4 gene located on chromosomal region 11q23. The age of onset is early. This disease is characterized by impairment of terminal steps of glycogenolysis and gluconeogenesis. Patients manifest a wide range of clinical symptoms and biochemical	600,25

Glycogen storage disease type 1B patients also present a tendency towards infections towards infections associated with neutropenia, relapsing aphthous gingivostomatitis, and inflammatory bowel disease. The incidence is 1:100,000. Congenital hereditary endothelial dystrophy type 2 follows an autosoma 2 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SLC4AII gene located on chromosomal region

20p13. The age of onset is 600,25 early. This disease is early. This disease is characterized by a diffuse ground-glass appearance of the corneas and marked corneal thickening from birth with nystagmus, and blurred vision

Spinal muscular atrophy follows an autosomal rollows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SMNI gene located on chromosomal region Sq13.2. The age of onset is variable. This disease. variable. This disease comprise a group of comprise a group of neuromuscular disorders characterized by degeneration of the anterior horn cells of the spinal cord, leading to symmetrical muscle weakness and atrophy weakness and atrophy.
Autosomal recessive forms
are classified according to
the age of onset, the
maximum muscular
activity achieved, and
survivorship. The severity of the disease is mainly of the disease is mainly determined by the copy number of SMN2, a copy gene which predominantly produces exon 7-skipped transcripts 600,25 and only low amount of full bearth transcripts has

and only low amount of full-length transcripts that encode for a protein identical to SMNI. Only about 4% of patients bear one SMNI copy with an intragenic mutation. Type 1 is a severe form, with onset before 6 months of onset perore 6 months or age. Patients never achieve the ability to sit. Type 2 has intermediate severity, with onset between 6 and 18 months.

Patients do not reach the motor milestone of

motor milestone of standing, and survive into adulthood. Type 3 onset is after 18 months. Patients develop ability to stand and walk and survive into adulthood. Type 4 onset is in adulthood, disease In adulthood, disease progression is slow, and patients can stand and walk. The incidence is 1:10,000 and the prevalence is 1:80,000. Niemann-Pick disease, type A and type B follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SMPD1 gene located on chromosomal region 11p15.4. The clinical phenotype ranges from a severe infantile form with

600.25

severe infantile form with neurologic degeneration resulting in death usually by 3 years of age (type A) to a later-onset nonneurologic form (type B) that is compatible with survival into adulthood. Since intermediate cases also have been reported, the disease is best regarded a single entity with a clinical spectrum.

Amyotrophic lateral sclerosis, type 5, juvenile follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the SPG11 gene located on chromosomal region 15q21.1. The age of onset is infancy/childhood. This disease is characterized by 600.25

progressive upper and lower motor neuron lower motor neuron degeneration causing facial spasticity, dysarthria, and gait disorders with onset before 25 years of age. The prevalence is <1/1,000,000.

Spastic paraplegia type 7 follows an autosomal recessive pattern of inheritance and is caused

NM_001174090.1c.2687G>A, NM_001174090.1c.2686C>T, NM_001174090.1c.2647A>G, NM_001174090.1c.2609T>C, NM_001174090.1c.265G>A, NM_001174090.1c.2314_2321dupTATCACAC, NM_001174090.1c.2305G>A, NM_001174090.1c.2305G>A, NM_001174090.1c.2305G>A, NM_001174090.1c.2305G>A, NM_001174090.1c.2015G>A, NM_001174090.1c.2015G>A, NM_001174090.1c.2015G>A, NM_001174090.1c.2015G>A, NM_001174090.1c.2015G>A, NM_001174090.1c.2015G>A, NM_001174090.1c.2015G>A, NM_001174090.1c.2015G-A, NM_001174 NM 001174090.1:c.1472G>A. NM 001174090.1:c.1119 1120insA NM_001174090.1:c.718T>C, NM_001174090.1:c.554_561delGCTTCGCC

NM_001174090.1

Corneal endothelial dystrophy, autosomal

SLC4A11

Spinal muscular atrophy -0

del ex7, del ex7-8

NM_000543.4c.96G-A, NM_000543.4c.103.107delCTGGT,
NM_000543.4c.36delG, NM_000543.4c.354delC,
NM_000543.4c.475T>C, NM_000543.4c.55TC>T,
NM_000543.4c.54delC, NM_000543.4c.55TC>T,
NM_000543.4c.57delC, NM_000543.4c.55ddupC,
NM_000543.4c.5730G-A, NM_000543.4c.588C>T,
NM_000543.4c.7370G-A, NM_000543.4c.742C>A,
NM_000543.4c.757G>C, NM_000543.4c.788T>A,
NM_000543.4c.75TG>C, NM_000543.4c.78ET>A,
NM_000543.4c.75TG>C, NM_000543.4c.75TC>C,
NM_000543.4c.75TC>T, NM_000543.4c.75TC>T,
NM_000543.4c.75TC, NM_000543.4c.75TC,
NM_000543.4c.75TC, NM_000543.4c.75TC,
NM_000543.4c.75TC, NM_000543.4c.75TC,
NM_000543.4c.75TC, NM_000543.4c.75TC,
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NM_000543.4c.75TC,
NM_000543.4c.75TC,
NM_00054

NM_000543.4:c.96G>A, NM_000543.4:c.103_107delCTGGT,

Niemann-Pick disease, SMPD1 NM 000543.4 type A and type B

Amyotrophic lateral sclerosis, type 5, juvenile SPG11

NM_025137.3

NM 025137.3:c.7152-1G>C. NM 025137.3:c.6847 6848dupTC. NM_025137.3c::7152-1G>C, NM_025137.3c::6847_6848dup1C, NM_025137.3c::6805_6806delCT, NM_025137.3c::6100C>T, NM_025137.3c::5623C>T, NM_025137.3c::1736-1G>C, NM_025137.3c::1339_1342dupGGCT, NM_025137.3c::733_734delAT, NM_025137.3c::342delT, NM_025137.3c::342delT, NM_025137.3c::18C>T

SPG7	Spastic paraplegia, type 7, autosomal recessive	NM_003119.3	NM_003119.3:c.233T>A, NM_003119.3:c.286+1G>T, NM_003119.3:c.679C> NM_003119.3:c.758+217>C, NM_003119.3:c.773_774de1TG, NM_003119.3:c.1045G>A, NM_003119.3:c.1124de1G, NM_003119.3:c.159C>T, NM_003119.3:c.1676de1A, NM_003119.3:c.1749G>C, NM_003119.3:c.2075G>C	the SPC7 gene located on chromosomal region 7, 16q24.3. The age of onset is adult. This disease is characterized by by progressive muscle stiffness (spasticity) and the development of paralysis of the lower limbs (paraplegia) due to degeneration of corticospinal axons. The prevalence is 1:100,000-9:100,000.	600,25
STRC	Deafness, autosomal recessive, type 16	NM_153700.2	NM_153700.2:c.5188C>T, NM_153700.2:c.5185C>T, NM_153700.2:c.5168_5171delTTCT, NM_153700.2:c.4560dupC, NM_153700.2:c.4545+1G>C, NM_153700.2:c.3556C>T	Autosomal recessive nonsyndromic sensorineural deafness type DFNB16 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the STRC gene located on chromosomal region 15q15.3. The age of onset is early. This disease is characterized by hearing loss and deafness, no associated visible abnormalities of the external ear or any related medical problems. Autosomal recessive limbgirdle muscular dystrophy	600,25
TCAP	Muscular dystrophy, limb- girdle, type 2G	NM_003673.3	NM_003673.3:c.157C>T	ype 2G follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TCAP gene located on chromosomal region 17912. The age of onset is variable. This disease is characterized by muscle weakness in the four limbs, mild scapular winging, severe atrophy of the quadriceps and anterior tibialis muscles, calf hypertrophy, and lack of respiratory and cardiac	600,25
TCIRGI	Osteopetrosis, autosomal recessive, type 1	NM_006019.3	NM_006019.3:c.115_116delGA, NM_006019.3:c.1213G>A, NM_006019.3:c.1331G>T, NM_006019.3:c.1674-1G>A, NM_006019.3:c.2236+1G>A	involvement. Autosomal recessive osteopetrosis type 1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TCIRG1 gene located on chromosomal region Ilq152. The age of onset is early. This disease is characterized by bone marrow failure, fractures and visual impairment. The incidence is 1:200.000 live births and the prevalence is 1:250,000.	600,25
TERT	Dyskeratosis congenita, autosomal recessive, type 4	NM_198253.2	NM_198253.2:e.2701C>T, NM_198253.2:e.2431C>T	Dyskeratosis congenita, autosomal recessive follows an autosomal recessive follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TERT gene located on chromosomal region 5p15.33. The age of onset is early. This disease has a wide phenotypic spectrum and age onset. It usually manifests during childhood with the triad of dysplastic nails, lacy reticular pigmentation and atrophy of the skin at the level of the neck and upper chest, and oral leukoplaskia. Patients show an increased risk for progressive bone marrow failure and may develop myelodysplastic syndrome or acute myelogenous leukemia at any age (the risk increasing with age). There is also an increased risk for solid tumors, typically squarmous cell carcinoma of head and neck (see this term) or anogenital cancer. Various additional clinical findings have been reported and may include: developmental delay, short stature, microcephaly, blepharitis, epiphora, periodontal disease, taurodontism, decreased teeth/root ratio, esophageal stenosis, liver disease, urethral stenosis, osteoporosis, avascular necrosis of femur and/or humerus, premature hair greying/alopecia, or abnormal eyelashes. Individuals with DC are at high risk of pulmonary fisions. The prevalence is 11,000,000.	600,25
TFR2	Hemochromatosis, type 3	NM_001206855.1	NM_001206855.1:c.2T>A	inheritance and is caused by pathogenic variants in the TFR2 gene located on chromosomal region 7q22.1. The age of onset is adult. This disease is characterized by excessive tissue iron deposition of genetic origin, liver	600,25

the SPG7 gene located on

TFR2	Hemochromatosis, type 3	NM_003227.3	NM_003227.3:c.2374G>A, NM_003227.3:c.2343G>A, NM_003227.3:c.2014C>T, NM_003227.3:c.1861_1872delGCCGTGGCCCAG, NM_003227.3:c.1655delC, NM_003227.3:c.1633delGA, NM_003227.3:c.147541G-A, NM_003227.3:c.147541G-A, NM_003227.3:c.1235_1237delACA, NM_003227.3:c.1330G>A, NM_003227.3:c.949C>T, NM_003227.3:c.186C>T, NM_003227.3:c.313C>T	aisease, nypogonadism, arthritis, diabetes and skin pigmentation. The prevalence is <1,100,000. Hemochromatosis type 3 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TFR2 gene located on chromosomal region 7q22.1. The age of onset is adult. This disease is characterized by excessive tissue iron deposition of genetic origin, liver disease, hypogonadism, arthritis, diabetes and skin pigmentation. The prevalence is <11,000,000. Mitochondrial DNA depletion syndrome type 2 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TK2 gene located on chromosomal region	600,25
TK2	Mitochondrial DNA depletion syndrome, type 2 (myopathic type)	NM_004614.4	NM_004614.4:c.635T>A, NM_004614.4:c.604_606delAAG, NM_004614.4:c.500G>A, NM_004614.4:c.373C>T, NM_004614.4:c.353C>A, NM_004614.4:c.323C>T, NM_004614.4:c.359C>G	IGQ1. The age of onset is infantile. This disease is characterized by generalized hypotonia, proximal muscle weakness, loss of previously acquired motor skills, poor feeding, and respiratory difficulties leading to respiratory failure and death within a few years after diagnosis. The prevalence is 1.2100,000. Joubert syndrome (JBTS) type 6 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TMEM67 gene located on chromosomal region 8,221. The age of onset is early, JBTS is characterized by congenital malformation of the brainstem and agenesis of the cerebellar vermis (molar tooth sign) leading to an abnormal respiratory pattern, nystagmus, hypotonia, mental retardation, ataxia, and delay in achieving motor milestones. Other variable features include retinal dystrophy (manifesting with either Leber congenital amaurosis or progressive retinal dystrophy) and	
тмем67	Joubert syndrome, type 6; Meckel syndrome, type 3; COACH syndrome	NM_153704.5	NM_153704.5:c.130C>T, NM_153704.5:c.14B_149insTAAT, NM_153704.5:c.622A>T, NM_153704.5:c.755T>C, NM_153704.5:c.1046T>C, NM_153704.5:c.1538A>C, NM_153704.5:c.1769T>C, NM_153704.5:c.2498T>C	nephronophthisis (usually	
TMPRSS3	Deafness, autosomal recessive, type 8/10	NM_024022.2	NM_024022.2:c.1276G>A, NM_024022.2:c.1211C>T, NM_024022.2:c.53G>C, NM_024022.2:c.647G>T, NM_024022.2:c.4646-16-T, NM_024022.2:c.43C>A, NM_024022.2:c.42C>G, NM_024022.2:c.20BdelC	sensorneural deatness type DFNBI0 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TMPRSS gene located on chromosomal region 21q22.3. The age of onset is early. This disease is characterized by hearing loss and deafness. Neuronal ceroid lipofuscinosis type 2 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TPPI gene located on chromosomal region	600,25
TPPI	Ceroid lipofuscinosis, neuronal, type 2	NM_000391.3	NM_000391.3:c.1340G>A, NM_000391.3:c.1093T>C, NM_000391.3:c.510>T, NM_000391.3:c.522C>T, NM_000391.3:c.509-1G>C, NM_000391.3:c.509-1G>C, NM_000391.3:c.141_144delGAGT	11p15.4. Age of onset is infantile. This disease is characterized by epilepsy,	600,25

TRIOBP	Deafness, autosomal recessive, type 28	NM_001039141.2	NM_001039141.2:c.1039C>T, NM_001039141.2:c.1741C>T, NM_001039141.2:c.2639_2640insTCAC, NM_001039141.2:c.3195delT, NM_001039141.2:c.3202C>T, NM_001039141.2:c.4577C>G, NM_001039141.2:c.5316G>A	rapidly progresses to light/dark awareness only. Life expectancy ranges from age six years to early teenage. The prevalence is 1.51,000,000-91,000,000. Deafness autosomal recessive type 28 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TRIOBP gene located on chromosomal region 22q13.1. The age of onset is early. This disease is characterized by hearing loss and deafness, no associated visible abnormalities of the external ear or any related medical problems. Pontocerebellar.),25
TSEN54	Pontocerebellar hypoplasia, type 2A	NM_207346.2	NM_207346.2:c.670_671delAA, NM_207346.2:c.736C>T, NM_207346.2:c.887G>A, NM_207346.2:c.919G>T, NM_207346.2:c.1027C>T, NM_207346.2:c.1039A>T	an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TSENS4 gene located on chromosomal region 17q2S1. Pontocerebellar hypoplasia (PCH) refers to a group of severe neurodegenerative disorders affecting growth and function of the brainstem and cerebellum, resulting in little or no development. Different types were classified based on the clinical picture and the spectrum of pathologic),25
TSFM	Combined oxidative phosphorylation deficiency, type 3	NM_001172696.1	NM_001172696.1:c.1_2delAT, NM_001172696.1:c.24_25delCG, NM_001172696.1:c.581delC, NM_001172696.1:c.919C>T	changes. Combined oxidative phosphorylation deficiency type 3 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TSFM gene located on chromosomal region 12q/41. The age of onset is early. This disease is characterized by hypotonia, lactic acidosis, and hepatic insufficiency, with progressive encephalomyopathy or hypertrophic),25
TSHR	Hypothyroidism, congenital, nongoitrous, type 1	NM_000369.2	NM_000369.2:c.122G>C, NM_000369.2:c.202C>T, NM_000369.2:c.484C>G, NM_000369.2:c.500T>A, NM_000369.2:c.1170T>G, NM_000369.2:c.1742dupC	cardiomyopathy. Hypothyroidism, congenital, nongoitrous, type1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TSHR gene located on chromosomal region 14q311. This disease is characterized by increased levels of plasma TSH and low levels of thyroid hormone. Only a subset of patients develop frank hypothyroidism; the remainder are euthyroid and asymptomatic.),25
TTN	Limb-girdle muscular dystrophy type 10 (LGMDR10; formerly LGMD21); Early-onset myopathy with fatal cardiomyopathy	NM_001267550.2	NM_001267550.2:c.107889delA, NM_001267550.2:c.106070_106071delAT NM_001267550.2:c.104092delC, NM_001267550.2:c.104092c>T, NM_001267550.2:c.98818_98821delTcCA, NM_001267550.2:c.603544c>G, NM_001267550.2:c.603681dupT, NM_001267550.2:c.69344c>G, NM_001267550.2:c.52372delG, NM_001267550.2:c.48253delA, NM_001267550.2:c.47915dupT, NM_001267550.2:c.48253delA, NM_001267550.2:c.47915dupT, NM_001267550.2:c.28300_28303delAGCA, NM_001267550.2:c.16881c>A, NM_001267550.2:c.15796c>T, NM_001267550.2:c.3165-1G>T	LGMDRIO is a severe recessive form of LGMD phenotype with onset in the first to third decades involving weakness of all proximal muscles. Severe disability with loss of ambulation may occur within 20 years (third to sixth decades). Most of the cases are without facial muscle involvement or cardiomyopathy. Some patients later developed distal muscle involvement. Early-onset myopathy with fatal cardiomyopathy also follows an autosomal recessive pattern of inheritance. This),25
ТТРА	Ataxia with isolated vitamin E deficiency	NM_000370.3	NM_000370.3:c.744delA, NM_000370.3:c.661C>T, NM_000370.3:c.575G>A	caused by pathogenic variants in the TTN gene located on chromosomal region 2q31.2. Ataxia with vitamin E deficiency follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TTPA gene located on chromosomal region 8q13. The age of onset is variable. This disease is characterized by progressive spinocerebellar ataxia, loss of proprioception, areflexia, and is associated with a marked deficiency in vitamin E. The prevalence is 0.561,000,000.),25

TYR	Albinism, oculocutaneous, type 1A	NM_000372.4	NM_000372.4:c.1A>G, NM_000372.4:c.140G>A, NM_000372.4:c.164G>A, NM_000372.4:c.23G>A, NM_000372.4:c.24C>T, NM_000372.4:c.25G>A, NM_000372.4:c.25G>A, NM_000372.4:c.25G>A, NM_000372.4:c.25G>A, NM_000372.4:c.325G>A, NM_000372.4:c.35G>A, NM_000372.4:c.35GO>A, NM_000372.4:c.35GO	type 1A follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TYR gene located on chromosomal	00,25
TYRPI	Albinism, oculocutaneous, type 3	NM_000550.2	NM_000550.2:c.107delT, NM_000550.2:c.176C>G, NM_000550.2:c.497C>G, NM_000550.2:c.1057_1060delAACA, NM_000550.2:c.1067G>A, NM_000550.2:c.1103delA, NM_000550.2:c.1120C>T, NM_000550.2:c.1372_1375dupGACA	autosomal recessive pattern of inheritance and is caused by pathogenic variants in the TYRPI gene located on chromosomal region 9p23. The age of onset is early. This disease is characterized by rufous or brown albinism and occurring mainly in the African population. The prevalence is of 1/8,500	00,25
UGTIAI	Crigler-Najjar syndrome, type 2	NM_000463.2	NM_000463.2:c.44T>G, NM_000463.2:c.1021C>T, NM_000463.2:c.1070A>G, NM_000463.2:c.1456T>G	individuals in Africa. Crigler-Najjar syndrome type 2 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the UCTIAI gene located on chromosomal region 2q37. The age of onset is early. This disease is characterized by unconjugated hyperbillirubinemia due to reduced and inducible activity of hepatic bilirubin glucuronosyltransferase with pigmented bile that contains bilirubin glucuronides, and generally do not present neurologic or intellectual impairment. Bilirubin encephalopathy may develop in later life when patients experience a superimposed infection or stress.	00.25
USHIC	Usher syndrome, type 1C; Deafness, autosomal recessive, type 18A	NM_153676.3	NM_153676.3:c.2688_2695dupAATTCACC, NM_153676.3:c.2622_2623delCA, NM_153676.3:c.2547-1G>T, NM_153676.3:c.238dupC, NM_153676.3:c.238delC, NM_153676.3:c.216G>A	Usher syndrome type IC follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the USHIC gene located on chromosomal region 1lp151. This disease is characterized by the association of sensorineural deafness (usually congenital, severe and stable), progressive vision loss caused by	00,25
USHZA	Usher syndrome, type 2A	NM_206933.2	NM_206933.2:c.15520-1G>A, NM_206933.2:c.15371delT, NM_206933.2:c.15890-A, NM_206933.2:c.14803C>T, NM_206933.2:c.15089C>A, NM_206933.2:c.13709delG, NM_206933.2:c.135709delG, NM_206933.2:c.13574C-T, NM_206933.2:c.1254_12255delGA, NM_206933.2:c.105615C-T, NM_206933.2:c.10563GC>A, NM_206933.2:c.1056175C, NM_206933.2:c.1056176C, NM_206933.2:c.1056176-T, NM_206933.2:c.1056176-T, NM_206933.2:c.5745_5774delAG, NM_206933.2:c.5745_5774delAG, NM_206933.2:c.5745_5774delAG, NM_206933.2:c.3991_3492delCT, NM_206933.2:c.289861G, NM_206933.2:c.289861G, NM_206933.2:c.289861G, NM_206933.2:c.289861G, NM_206933.2:c.289661G, NM_206933.2:c.28961G, NM_206933.2:c.8969361G, NM_206933.2:c.89661G, NM_206933.2:c.99661G, NM_2	recessive pattern of inheritance and is caused by pathogenic variants in the USH2A gene located on chromosomal region 1q41. This disease is characterized by the association of sensorineural deafness (usually congenital, moderate/severe and stable) and progressive vision loss that begins in adolescence or adulthood caused by retinitis pigmentosa. Unlike the other forms of Usher syndrome, type 2 is not associated with vestibular abnormalities that cause difficulties with balance. USH2A accounts for more than half of all cases of Usher syndrome type 2. The estimated prevalence	00.25
WFSI	Wolfram syndrome, type 1	NM_001145853.1	NM_001145853.1c.616C>T, NM_001145853.1c.676C>T, NM_001145853.1c.1060_1062delTTC, NM_001145853.1c.1050_1052delTTC, NM_001145853.1c.1233delCTCT, NM_001145853.1c.1234_1237delGTCT, NM_001145853.1c.1511C>T, NM_001145853.1c.234_1237delGTCT, NM_001145853.1c.1944G>A, NM_001145853.1c.2643_2644delCT	is 3:100,000-4:100,000. Wolfram syndrome, type 1 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the WFSI gene located on chromosomal region 4p16.1. The age of onset is infantile. This disease is characterized by diabetes mellitus type I, diabetes insipidus, optical atrophy and neurological signs. The prevalence is 1:1,000,000-9:1,000,000. Usher syndrome type 2D	00.25

WHRN	Usher syndrome, type 2D; Deafness, autosomal recessive, type 31	NM_015404.3	NM_015404.3:c.817C>T	follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the WHRN gene located on chromosomal region 9q32. This disease is characterized by the association of sensorineural deafness (usually congenital, moderate/severe and stable) and progressive vision loss that begins in adolescence or adulthood caused by retinitis pigmentosa. Unlike the other forms of Usher syndrome, type 2 is not associated with vestibular abnormalities that cause difficulties with balance. The WHRN gene is also associated with autosomal recessive nonsyndromic sensorineural deafness type 31. This phenotype is characterized by congenital, non-progressive, mild-to-profound sensorineural	600,25
WNTIOA	Odontoonychodermal dysplasia	NM_025216.2	NM_025216.2:c.321C>A, NM_025216.2:c.383G>A, NM_025216.2:c.697G>T	hearing impairment. Odonto-onycho-dermal dysplasia follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the WNTIOA gene located on chromosomal region 2q35. The age of onset is infantile. This disease is characterized by hyperkeratosis and hyperhidrosis of the palms and soles, atrophic malar patches, hypodontia, conical teeth, onychodysplasia, and dry and sparse hair. The prevalence is <11,000,000.	600,25
ZFVVE26	Spastic paraplegia, type 15, autosomal recessive	NM_015346.3	NM_015346.3:c.5485-1G>A, NM_015346.3:c.5422C>T, NM_015346.3:c.4936C>T, NM_015346.3:c.4312C>T, NM_015346.3:c.3205C>T, NM_015346.3:c.3206G>A, NM_015346.3:c.3182delT, NM_015346.3:c.314dupC, NM_015346.3:c.1477C>T	Spastic paraplegia type 15 follows an autosomal recessive pattern of inheritance and is caused by pathogenic variants in the ZFYVE26 gene located on chromosomal region 14q24.1. The age of onset is infancy. This disease is characterized by progressive spasticity primarily affecting the lower limbs. It is a complex form of spastic paraplegia, associated with other neurologic dysfunction, including variable mental retardation, hearing and visual defects, and thin corpus callosum. The prevalence is <1/	600,25