(2021-03-11) רשימת מחלות ומוטציות בפאנל Clalit Hybrid Capture-Based Next Generation Sequencing

Mutation #	Gene: Disease Name {(Transcsript) "Mutation name"}
1	2p21:Hypotonia-cystinuria syndrome {"2p21"}
2 - 6	ABCA12:Ichthyosis, congenital, Autosomal Recessive 4A {(NM_173076.3) "c.1060C>T", "c.179G>C", "c.3456G>A", "c.4544G>A", "c.4553G>A"}
7 - 7	ABCA3:Congenital surfactant deficiency (Surfactant metabolism dysfunction, pulmonary, 3) {(NM_001089) "c.1474dupT"}
8 - 13	ABCA4:Cone-rod dystrophy 3 {(NM_000350.3) "c.1648G>A", "c.2791G>T", "c.3607G>A", "c.3608G>A", "c.5460+1G>A", "c.834delT"}
14 - 16	ABCB11:Cholestasis, progressive familial intrahepatic type 2 {(NM_003742) "c.1100_1101insTA" (NM_003742.4) "c.1409G>A", "c.3268C>T"}
17 - 22	ABCC8:Hyperinsulinemic hypoglycemia, familial, 1 (PHHI/HHF1) {(NM_000352) "c.3339dupG" (NM_000352.4) "c.1116dupT", "c.2506C>T", "c.3989-9G>A", "c.4160_4162delTCT", "c.560T>A"}
23 - 23	ABCD1:X-linked adrenoleukodystrophy {(NM_000033.4) "c.686T>C"}
24 - 25	ABHD5:Chanarin-Dorfman syndrome {(NM_016006.6) "c.412T>C", "c.934C>T"}
26 - 36	ACADM:Medium-chain Acyl-CoA dehydrogenase deficiency {(NM_000016.5) "c.1010A>C", "c.1045C>T", "c.1257C>A", "c.362C>T", "c.415_419delGATCA", "c.431_434delAGTA", "c.454G>T", "c.616C>T", "c.621_624delTGAT", "c.799G>A", "c.985A>G"}
37 - 45	ACADVL:Acyl-CoA dehydrogenase, very long-chain, VLCAD deficiency {(NM_000018.4) "c.1096C>T", "c.1748C>T", "c.260T>C", "c.367G>A", "c.637G>A", "c.65C>A", "c.779C>T", "c.799_802delGTTA", "c.894G>A"}
46 - 46	ACO2:Infantile cerebellar-retinal degeneration {(NM_001098.3) "c.336C>G"}
47 - 48	ACP5:Spondyloenchondrodysplasia with immune dysregulation {(NM_001111035) "c.772_790del" (NM_001111035.2) "c.325G>A"}
49 - 49	ACSF3:Combined malonic and methylmalonic aciduria {(NM_174917) "c.1411C>T"}
50 - 51	ADA:Severe combined immunodeficiency due to ADA deficiency {(NM_000022.4) "c.703C>T", "c.792G>A"}
52 - 52	ADAM9:cone-rod dystrophy 9 {(NM_003816) "c.1087T>A"}
53 - 54	ADAMTS2:Ehlers Danlos syndrome, type VIIC {(NM_014244.5) "c.2384G>A", "c.673C>T"}
55 - 59	ADGRG1:Bilateral Frontoparietal Polymicrogyria (BFPP) {(NM_005682.7) "c.1036T>A", "c.1046G>C", "c.1167+3G>C", "c.1693C>T", "c.739_745delCAGGACC"}
60 - 61	ADGRV1:Usher syndrome, type 2C {(NM_032119.4) "c.14973-2A>G", "c.15494delA"}
62 - 62	AGA:Aspartylglucosaminuria {(NM_000027.4) "c.214T>C"}
63 - 67	AGL:Glycogen storage disease III {(NM_000642.3) "c.1078C>T", "c.1222C>T", "c.2812+2dupT", "c.3652C>T", "c.4456delT"}

68 - 82	AGXT:Hyperoxaluria, primary, type I {(NM_000030) "c.26C>A" (NM_000030.3) "c.121G>A", "c.33delC", "c.33dupC", "c.466G>A", "c.584T>G", "c.586G>A", "c.680+1G>A", "c.697C>T", "c.727G>C",
	"c.731T>C", "c.860_861delGCinsCG", "c.865C>T", "c.893T>C", "c.997A>T"}
83 - 85	AHI1:Joubert syndrome-3 {(NM_017651.4) "c.2212C>T", "c.3032C>G", "c.787dupC"}
86 - 86	AIMP1:Leukodystrophy, hypomyelinating, 3 {(NM_004757.3) "c.292_293delCA"}
87 - 89	AIPL1:Leber congenital amaurosis 4 {(NM_014336.5) "c.211G>T", "c.215G>A", "c.834G>A"}
90 - 95	AIRE:Autoimmune polyendocrinopathy syndrome, type I (APS-1) with or without reversible metaphyseal dysplasia {(NM_000383.4) "c.1163_1164insA", "c.247A>G", "c.254A>G", "c.44G>A", "c.47C>T", "c.769C>T"}
96 - 96	ALDH1A3:Microphthalmia, isolated 8 {(NM_000693.4) "c.211G>A"}
97 - 97	ALDH7A1:Epilepsy, pyridoxine-dependent {(NM_001182.5) "c.1489+5G>A"}
98 - 105	ALDOB:Fructose intolerance {(NM_000035) "c.178C>T", "c.360_363delCAAA", "c.612T>A", "c.612T>G", "c.865delC" (NM_000035.4) "c.1005C>G", "c.448G>C", "c.524C>A"}
106 - 108	ALMS1:Alstrom syndrome {(NM_015120.4) "c.8008C>T", "c.808C>T", "c.8171_8181del"}
109 - 111	ALPL:Hypophosphatasia, infantile {(NM_000478) "c.1337delC" (NM_000478.6) "c.1348C>T", "c.141C>A"}
112 - 112	AMT:Glycine encephalopathy, AMT-related {(NM_000481.3) "c.125A>G"}
113 - 113	ANO5:Limb-girdle muscular dystrophy {(NM_213599.2) "c.191dupA"}
114 - 114	AP4B1:Spastic paraplegia 47, Autosomal Recessive {(NM_006594.4) "c.664delC"}
115 - 115	APTX:Ataxia, early-onset, with oculomotor apraxia and hypoalbuminemia {(NM_175073.2) "c.837G>A"}
116 - 117	AQP2:Diabetes insipidus, nephrogenic {(NM_000486.5) "c.298G>T", "c.83T>C"}
118 - 118	ARFGEF2:Periventricular heterotopia with microcephaly {(NM_006420.3) "c.1958+1G>A"}
119 - 119	ARHGDIA:Nephrotic syndrome, type 8 {(NM_004309.6) "c.518G>T"}
120 - 120	ARL6:Bardet-Biedl syndrome 3 {(NM_032146.5) "c.364C>T"}
121 - 134	ARSA:Metachromatic leukodystrophy - MLD {(NM_000487)
	"c.1114C>T" (NM_000487.6) "c.1136C>T", "c.1174C>T",
	"c.1283C>T", "c.211T>G", "c.263G>A", "c.292_293delTCinsCT",
	"c.937C>T" (NM_001085425.3) "c.449C>T"}
135 - 135	ARSG:Usher syndrome, type IV {(NM_014960) "c.133G>T"}
136 - 136	ASL:Argininosuccinic aciduria {(NM_000048.4) "c.346C>T"}
137 - 137	ASNS:Asparagine synthetase deficiency {(NM_183356.3) "c.1084T>G"}
138 - 141	ASPA:Canavan Disease {(NM_001128085.1) "c.433-2A>G",

	"c.693C>A", "c.854A>C", "c.914C>A"}
142 - 142	ASS1:Citrullinemia,classic {(NM_000050) "c.1168G>A"}
143 - 156	ATM:Ataxia-tyelangiectasia {(NM_000051) "c.1514T>C" (NM_000051.3) "c.103C>T", "c.1339C>T", "c.1547T>C", "c.2284_2285delCT", "c.2839-579_2839-576del4", "c.3245_3247delATCinsTGAT", "c.3576G>A", "c.368delA", "c.497del7514", "c.5763-1050A>G",
	"c.6672_6680delGGCTCTACGinsCTC", "c.7241_7244delAAGC", "deletion_exon_3-4"}
157 - 157	ATP6V0A2:Cutis laxa, Autosomal Recessive, type IIA {(NM_012463.4) "c.2375C>G"}
158 - 180	ATP7B:Wilson disease {(NM_000053) "c.3007G>A", "c.3784G>T" (NM_000053.3) "c.122A>G", "c.1340_1343delAAAC", "c.1544G>A", "c.1639delC", "c.1703T>G", "c.1934T>G", "c.2293G>A", "c.2333G>T", "c.2337G>A", "c.2817G>T", "c.2906G>A", "c.3191A>C", "c.3207C>A", "c.3451C>T", "c.3551T>C", "c.3638G>T", "c.3649_3654delGTTCTG", "c.3659C>T", "c.3842G>A", "c.4152T>G", "c.845delT"}
181 - 181	ATP8B1:Cholestasis, progressive familial intrahepatic 1 {(NM_005603) "c.2854C>T"}
182 - 182	AVP:Familial neurohypophyseal diabetes insipidus {(NM_000490) "c.77C>T"}
183 - 183	B3GALNT2:Muscular dystrophy-dystroglycanopathy (congenital with brain and eye anomalies, type A, 11) {(NM_152490.5) "c.236-1G>C"}
184 - 184	B4GALT1:Congenital disorder of glycosylation, type IId {(NM_001497) "c.61C>T"}
185 - 187	BBS10:Bardet-Biedl syndrome 10 {(NM_024685.4) "c.1091delA", "c.1399delA", "c.271dupT"}
188 - 189	BBS1:Bardet-Biedl syndrome 1 {(NM_024649.5) "c.1169T>G", "c.479G>A"}
190 - 194	BBS2:Bardet-Biedl syndrome 2 {(NM_031885.4) "c.1895G>C", "c.224T>G", "c.311A>C", "c.401C>G", "c.98C>A"}
195 - 196	BBS4:Bardet-Biedl syndrome 4 {(NM_033028.5) "c.77-1421_221-1229del5784", "c.884G>C"}
197 - 197	BBS7:Bardet-Biedl syndrome 7 {(NM_176824.3) "c.1786G>A"}
198 - 199	BBS9:Bardet-Biedl syndrome 9 {(NM_014451) "c.1063C>T", "c.1669+1G>A"}
200 - 206	BCKDHA:Maple syrup urine disease, type Ia {(NM_000709.4) "c.169delG", "c.718del", "c.792C>G", "c.859C>T", "c.890G>A", "c.909_910delGT", "c.935_937del"}
207 - 213	BCKDHB:Maple syrup urine disease, type Ib {(NM_000056.4) "c.1016C>T", "c.1114G>T", "c.356T>G", "c.548G>C", "c.670C>T", "c.800_803delAGGA", "c.832G>A"}
214 - 220	BLM:Bloom syndrome {(NM_000057) "c.1642C>T", "c.2512C>T" (NM_000057.4) "c.1984_1985delAA", "c.2207_2212delATCTGAinsTAGATTC", "c.2407dupT", "c.3510T>A", "c.98+1G>T"}
221 - 221	BMPER:Diaphanospondylodysostosis {(NM_133468.5) "c.310C>T"}
222 - 222	BMPR1B:Brachydactyly type A2 {(NM_001256793.2) "c.377G>A"}

223 - 224	BSND:Bartter syndrome, type 4a infantile variant with sensorineuronal deafness {(NM_057176.3) "c.167_168insTTTCCC", "c.28G>A"}
225 226	,
225 - 226	BTD:Biotinidase deficiency {(NM_000060) "c.393delC" (NM_000060.4) "c.100G>A"}
227 - 229	C12ORF65:Spastic paraplegia 55, Autosomal Recessive {(NM_152269) "c.346delG" (NM_152269.5) "c.282+2T>A", "c.413_417delAACAA"}
230 - 230	C21orf59:Ciliary dyskinesia, primary, 26 {(NM_021254.4) "c.735C>G"}
231 - 236	C2ORF71:Retinitis pigmentosa 54 {(NM_001029883) "c.2334T>A",
	"c.2756_2768delAGCCAGCCCTGGA", "c.3289C>T", "c.478_479insA", "c.556C>T", "c.776_777delAG"}
237 - 239	C8orf37:Retinitis pigmentosa 64 {(NM_177965.4) "c.497T>A",
	"c.529C>T", "c.545A>G"}
240 - 242	CAPN3:Muscular dystrophy, limb-girdle, type 2A {(NM_000070) "c.1076C>T", "c.1469G>A", "c.367C>A"}
243 - 243	CASQ2:Ventricular tachycardia, catecholaminergic polymorphic, 2 {(NM_001232.3) "c.919G>C"}
244 - 248	CBS:Homocystinuria, thrombosis, hyperhomocysteinemic
	{(NM_000071.2) "c.1006C>T", "c.1224-2A>C", "c.1261delG",
	"c.785C>G" (NM_001178008.2) "c.919G>A"}
249 - 249	CC2D1A:Mental retardation, Autosomal Recessive 3 - MRT3
	{(NM_017721.5) "c.1468+1_1824-1del"}
250 - 250	CC2D2A:Mental retardation, Autosomal Recessive 3 - MRT3
	{(NM_001080522) "c.308delG"}
251 - 251	CCDC114:Ciliary dyskinesia, primary, 20 {(NM_144577) "c.939delT"}
252 - 252	CCDC174:Birk Volodarsky PMR Synderome Hypotonia and
	psychomotor developmental delay {(NM_016474.5) "c.1404A>G"}
253 - 253	CCDC65:Ciliary dyskinesia, primary, 27 {(NM_033124.5)
	"c.877_878delAT"}
254 - 254	CCDC88C:Hydrocephalus, nonsyndromic, Autosomal Recessive
	{(NM_001080414.4) "c.934C>T"}
255 - 258	CCNO:Ciliary dyskinesia, primary, 29 {(NM_021147) "c.165delC",
	"c.258_262dupGGCCC", "c.481_482delCT", "c.638T>C"}
	CD59:Hemolytic anemia & immune-mediated polyneuropathy, CD59-
1	related {(NM_203330.2) "c.266G>A"}
260 - 260	CDAN1:Dyserythropoietic anemia, congenital, type Ia
	{(NM_138477.4) "c.3124C>T"}
	CDH23:Usher Syndrome Type ID {(NM_022124.6) "c.7903G>T"}
	CDK10:Al Kaissi syndrome {(NM_052988) "c.870_871del"}
263 - 263	CDK5:Lissencephaly 7 with cerebellar hypoplasia {(NM_004935.4) "c.580+1G>A"}
264 - 264	CEACAM16:Deafness, autosomal recessive {(NM_001039213) "c.703C>T"}
265 - 265	CECR1:Adenosine deaminase 2 deficency {(NM_001282225.2)
	"c.140_141insT"}
266 - 266	CECR1:Polyarteritis nodosa, childhood-onset {(NM_001282225.2)
	"c.139G>A"}

267 - 267	CENPJ:Microcephaly, primary, Autosomal Recessive {(NM_018451.5) "c.3243_3246delTCAG"}
268 - 268	CEP104:Joubert syndrome (JBTS) {(NM_014704.4)
269 - 269	CEP152:Microcephaly 9, primary, Autosomal Recessive {(NM_014985.3) "c.2281-2A>G"}
270 - 278	CEP290:Meckel syndrome 4 {(NM_025114) "c.5668G>T",
	"c.5824C>T", "c.6760A>T" (NM_025114.3) "c.1225delA",
	"c.164_167delCTCA", "c.1666delA", "c.4393C>T", "c.4771C>T", "c.5788A>T"}
279 - 279	CERKL:Retinitis pigmentosa 26 {(NM_001030311.2) "c.238+1G>A"}
280 - 280	CFH:Hemolytic uremic syndrome, complement factor H deficiency
	{(NM_000186.3) "c.3677_*4del and c.3674A>T"}
281 - 708	CFTR:Cystic fibrosis {(NM_000492) "c.220C>T", "c.54-
	5811_164+2186del8108ins182" (NM_000492.3) "c.1000C>T",
	"c.1001G>A", "c.1007T>A", "c.1013C>T", "c.1021T>C",
	"c.1021_1022dupTC", "c.1040G>A", "c.1040G>C", "c.1040G>T",
	"c.1055G>A", "c.1075C>A", "c.1081delT", "c.1116+1G>A",
	"c.1130dupA", "c.1155_1156dupTA", "c.115C>T", "c.11C>A", "c.1202G>A", "c.1203G>A", "c.1209+1G>A", "c.1240C>T",
	"c.1327_1330dupGATA", "c.1340delA", "c.1364C>A", "c.137C>A",
	"c.1393-1G>A", "c.1397C>A", "c.1397C>G", "c.1400T>C", "c.1418delG",
	"c.1438G>T", "c.1439G>A", "c.1466C>A", "c.1475C>T", "c.1477C>T",
	"c.1521_1523delCTT", "c.1545_1546delTA", "c.1558G>A",
	"c.1558G>T", "c.1572C>A", "c.1573C>T", "c.1585-1G>A", "c.1585-
	8G>A", "c.1624G>T", "c.1625G>A", "c.1645A>C", "c.1646G>A",
	"c.1646G>T", "c.1647T>G", "c.165-1G>A", "c.1650delA", "c.1651G>A",
	"c.1652G>A", "c.1654C>T", "c.1657C>T", "c.166G>A", "c.1673T>C",
	"c.1675G>A", "c.1679+1634A>G", "c.1679+1G>C", "c.1679G>A",
	"c.1679G>C", "c.1680-1G>A", "c.1682C>A", "c.1692delA",
	"c.1736A>G", "c.1753G>T", "c.175dupA", "c.1766+1G>A",
	"c.1766+1G>C", "c.1766+3A>G", "c.1766+5G>T", "c.178G>T",
	"c.1817_1900del84", "c.1841A>G", "c.1911delG",
	"c.1923_1931delCTCAAAACTinsA",
	"c.1973_1985delGAAATTCAATCCTinsAGAAA", "c.1976delA",
	"c.1986_1989delAACT", "c.1A>G", "c.200C>T", "c.2012delT", "c.2051_2052delAAinsG", "c.2052_2053insA", "c.2052delA",
	"c.2089dupA", "c.2125C>T", "c.2128A>T", "c.2175dupA",
	"c.2195T>G", "c.2215delG", "c.223C>T", "c.2290C>T", "c.233dupT",
	"c.2353C>T", "c.2374C>T", "c.2423_2424dupAT", "c.2453delT",
	"c.2463_2464delTG", "c.2464G>T", "c.2490+1G>A", "c.2491G>T",
	"c.2537G>A", "c.2547C>A", "c.254G>A", "c.2551C>T", "c.2583delT",
	"c.2619+1G>A", "c.2619+2dupT", "c.262_263delTT",
	"c.2657+2_2657+3insA", "c.2657+5G>A", "c.2658-1G>C ",
	"c.2668C>T", "c.273+1G>A", "c.273+3A>C", "c.2737_2738insG",
	"c.2739T>A", "c.274-1G>A", "c.274G>A", "c.274G>T",
	"c.2763_2764dupAG", "c.2780T>C", "c.2834C>T", "c.2856G>C",
	"c.2875delG", "c.2908G>C", "c.292C>T", "c.2930C>T",
	"c.2988+1173_c.3468+2111del8898", "c.2988+1G>A", "c.2988G>A",

"c.2989-1G>A", "c.2989-977_3367+248del", "c.3002_3003delTG", "c.3039delC", "c.3067_3072delATAGTG", "c.3139+10T>C", "c.313delA", "c.3140-26A>G", "c.3154T>G", "c.3160C>G", "c.3181G>C", "c.3194T>C", "c.3196C>T", "c.3197G>A", "c.3205G>A", "c.3208C>T", "c.3209G>A", "c.3222T>A", "c.3230T>C" "c.325 327delTATinsG", "c.3266G>A", "c.3276C>G", "c.328G>C", "c.3292T>C", "c.3299A>C", "c.3302T>A", "c.3310G>T", "c.343G>T", "c.3454G>C", "c.3469-2A>G", "c.3472C>T", "c.3484C>T", "c.349C>T", "c.350G>A", "c.3528delC", "c.3529A>T", "c.3532_3535dupTCAA", "c.3535_3538delACCA", "c.3587C>G", "c.3600delA", "c.3605delA" "c.3611G>A", "c.3612G>A", "c.3659delC", "c.366T>A", "c.3691delT ", "c.3700A>G", "c.3712C>T", "c.3718-1G>A", "c.3718-2477C>T", "c.3731G>A", "c.3744delA", "c.3752G>A", "c.3763T>C", "c.3764C>A", "c.3764C>T", "c.3773dupT", "c.3793G>A", "c.3808delG", "c.3846G>A", "c.3873+1G>A", "c.3883_3884insG", "c.3883_3886delATTT", "c.3883delA", "c.3889dupT", "c.3909C>G", "c.3937C>T", "c.3964-78_4242+577del", "c.4046G>A", "c.4077_4080delTGTTinsAA", "c.4111G>T", "c.413 415dupTAC", "c.416A>C", "c.416A>T", "c.4197_4198delCT", "c.422C>A", "c.4234C>T", "c.4242+1G>T", "c.4251delA", "c.4297G>A", "c.4300_4301dupAG", "c.4364C>G", "c.442delA", "c.487A>G", "c.489+1G>T", "c.523A>G", "c.531delT" "c.532G>A", "c.54-5940 273+10250del21Kb", "c.543 546delTAGT", "c.575A>G", "c.579+1G>T", "c.579+3A>G", "c.579+5G>A", "c.580-1G>T", "c.595C>T", "c.613C>T", "c.617T>G", "c.658C>T", "c.675T>A", "c.761delA", "c.803delA", "c.805_806delAT", "c.828C>A", "c.870-2A>G", "c.892delA", "c.933_935delCTT", "c.948delT", "c.988G>T", "c.[1075C>A;1079C>A]" | (NM_000492.4) "c.-9_14del23", "c.1001G>T", "c.1006_1007insG", "c.1029delC", "c.1037T>C", "c.1046C>T", "c.1054C>T", "c.1079C>A", "c.1117-1G>A", "c.1211delG", "c.1301_1307delCACTTCT", "c.1327G>T", "c.1358T>C", "c.1365_1366delGG", "c.1367T>C", "c.1373delG", "c.1393-2A>G", "c.1408G>C", "c.1408G>T", "c.1420G>A", "c.1477_1478delCA", "c.1487G>A", "c.14C>T", "c.1505T>C", "c.1519_1521delATC", "c.1538A>G", "c.1584+1G>A", "c.164+1G>A", "c.164+1G>T", "c.164+2T>C", "c.164+3_164+4insT", "c.1648G>T", "c.165-3C>T", "c.1670delC", "c.1679+1G>A", "c.1680-877G>T", "c.1680A>C", "c.1687T>A", "c.1687T>G", "c.169T>G", "c.1703delT", "c.1705T>G", "c.170G>A or c.171G>A", "c.1721C>A", "c.1724T>A", "c.174_177delTAGA", "c.1763A>T", "c.1766+1G>T", "c.178G>A", "c.1792_1798delAAAACTA", "c.1801A>T", "c.1826A>G", "c.1837G>A", "c.1853T>C", "c.1865G>A", "c.1882G>C or c.1882G>A", "c.1923_1931del9insA", "c.1943delA", "c.1973_1985del13insAGAAA", "c.2017G>T", "c.2053C>T", "c.2053dupC", "c.2143C>T", "c.2158C>T", "c.2241_2248delGATACTGC", "c.2249C>T", "c.2537G>A or c.2538G>A", "c.2589_2599delAATTTGGTGCT", "c.2601dupA", "c.263T>A", "c.263T>G", "c.2645G>A", "c.271G>A", "c.274-2A>G", "c.2770G>A", "c.2810dupT", "c.2825delT", "c.2855T>C", "c.2859 2890delACATTCTGTTCTTCAAGCACCTATGTCAACCC", "c.2896delA", "c.2900T>C", "c.2909G>A", "c.2936A>T", "c.296C>T", "c.2989-2A>G", "c.3011_3019delCTATAGCAG or

	c.3009_3017delAGCTATAGC", "c.3017C>A", "c.3039dupC", "c.3047T>C", "c.305T>G", "c.3095A>G", "c.3107C>A", "c.310delA", "c.3124C>T", "c.3139_3139+1delGG", "c.3217dupT", "c.3220T>C", "c.3222T>G", "c.3276C>A", "c.3293G>A or c.3294G>A", "c.3294G>C or c.3294G>T", "c.3297C>A", "c.3302T>G", "c.3304A>T", "c.330C>A", "c.3468+2dupT", "c.3468+5G>A", "c.34686>A", "c.3475T>C", "c.3468+2dupT", "c.3468+5G>A", "c.3468G>A", "c.3475T>C", "c.3476C>T", "c.349C>G", "c.350G>C", "c.350G>T", "c.358G>A", "c.3717+40A>G", "c.3717+4A>G", "c.3717-5G>A", "c.3718-3T>G", "c.3719T>G", "c.3737C>T", "c.3806T>A", "c.3848G>T", "c.3872A>G", "c.3761T>G", "c.3737G>A", "c.3891dupT", "c.38C>T", "c.3908delA", "c.3971T>C", "c.3988C>T", "c.4004T>C", "c.4036_4042del", "c.4086dupT", "c.4097T>A", "c.409delC", "c.4124A>C", "c.4127_4131delTGGAT", "c.4144C>T", "c.4147dupA", "c.416A>G", "c.4231C>T", "c.4242+1G>A", "c.4426C>T", "c.44T>C", "c.470_483delTTAGTTTGATTTAT", "c.481T>G", "c.489+3A>G", "c.577G>T", "c.57G>A", "c.580G>A", "c.581G>T", "c.601G>A", "c.647G>A", "c.576SA", "c.580G>A", "c.79G>T", "c.601G>A", "c.647G>A", "c.680T>G", "c.695T>A", "c.79G>T", "c.825C>G", "c.772A>G", "c.794T>G", "c.79G>T", "c.88C>T", "c.88C>T", "c.88C>T", "c.8933C>G", "c.931G>A", "c.987delA"}
709 - 712	CHRNE:Myasthenic syndrome, congenital, 4B, fast-channel {(NM_000080) "c.1161_1162insT", "c.1353dupG", "c.187_188insC", "c.637dupG"}
713 - 718	CLCN1:Myotonia congenita, Autosomal Recessive {(NM_000083) "c.1012C>T", "c.1437_1450del", "c.1444G>A", "c.1586C>T", "c.568_569delGGinsTC", "c.803C>T"}
719 - 724	CLCN5:Proteinuria, low molecular weight, with hypercalciuric nephrocalcinosis {(NM_000084.4) "c.1245delG", "c.1399C>T", "c.161dup", "c.1909C>T", "c.258delA", "c.82C>T"}
725 - 726	CLCNKB:Bartter syndrome, type 3 and Gitelman syndrome {(NM_000085) "c.1830G>A" (NM_000085.4) "c.1313G>A"}
727 - 727	CLN5:Ceroid lipofuscinosis, neuronal, 5 {(NM_006493.3) "c.672delG"}
728 - 729	CLN6:Ceroid lipofuscinosis, neuronal, 6 {(NM_017882) "c.843G>A" (NM_017882.3) "c.214G>T"}
730 - 730	CLN8:Neuronal ceroid lipofuscinosis type 8, including northern epilepsy {(NM_018941.3) "c.766C>G"}
731 - 734	CLRN1:Usher syndrome, type 3A {(NM_174878.2) "c.144T>G", "c.349_358del", "c.433+1G>A", "c.528T>G"}
735 - 736	CNGA1:Retinitis pigmentosa 49 {(NM_000087) "c.1540C>T", "c.94C>T"}
737 - 745	CNGA3:Achromatopsia-2 - total color blindness {(NM_001298)
746 - 747	CNGB1:Retinitis pigmentosa 45 {(NM_001297) "c.2760G>A" (NM_001297.5) "c.2284C>T"}

748 - 755	CNGB3:Achromatopsia-3,macular degeneration, juvenile
	{(NM_019098) "c.1207C>T", "c.2328delC", "c.41_42dupTA",
	"c.467C>T", "c.819delC" (NM_019098.4) "c.1006G>T", "c.1148delC",
	"c.644-1G>C"}
756 - 757	CNNM4:Jalili syndrome {(NM_020184.4) "c.1813C>T", "c.599C>A"}
758 - 758	CNTNAP1:Lethal congenital contracture syndrome 7 {(NM_003632)
	"c.2015G>A"}
759 - 759	COL11A2:Otospondylomegaepiphyseal dysplasia (ZW)
	{(NM_080680.2) "c.3991C>T"}
760 - 764	COL17A1:Epidermolysis bullosa, junctional, {(NM_000494)
700 704	"c.2226insTGGA", "c.3676C>T", "c.3766+1G>A",
	"c.4145_4148delAGAG", "c.737_738insA"}
765 - 768	COL4A3:Alport Syndrome, COL4A3-Related {(NM_000091.4)
703 - 700	"c.1791_1793dupTCC", "c.227delG", "c.3518-2A>G", "c.4649T>G"}
769 - 770	COL4A4:Alport syndrome, COL4A4-Related {(NM 000092.4)
703-770	"c.3933C>G", "c.785_792dupCACCTGAC"}
771 - 776	COL4A5:Alport syndrome, COL4A5-Related {(NM_000495)
//1-//0	"c.1571delG" (NM_000495.4) "c.2641G>T", "c.367delG",
	"c.4691G>C", "c.4946T>G", "c.5030G>A"}
777 - 780	COL7A1:Dystrophic epidermolysis bullosa, Autosomal
/// - /80	,
	Recessive,COL7A1-Related {(NM_000094) "c.2387G>A", "c.4888C>T", "c.6341delG", "c.682+1G>A"}
704 704	
781 - 781	COLEC11:3MC syndrome 2 {(NM_199235.2) "c.627_628delCG"}
782 - 787	COLQ:Myasthenic syndrome, congenital, 5 {(NM_005677)
	"c.377delG", "c.788dupC", "c.893delA" (NM_005677.4) "c.1228C>T",
700 700	"c.718G>T", "c.792dupG"}
788 - 788	COQ4:Coenzyme Q10 deficiency, primary, 7 {(NM_016035.5) "c.718C>T"}
700 704	,
789 - 794	CPS1:Carbamoylphosphate synthetase I deficiency {(NM_001875.5)
	"c.1760G>A", "c.3265C>T", "c.3374C>T", "c.3558+1G>C",
705 705	"c.4101+2T>C", "c.794C>T"}
795 - 795	CPT1A:Carnitine palmitoyltransferase 1 deficiency {(NM_001031847) "c.1361A>G"}
706 707	,
796 - 797	CPT2:CPT deficiency, hepatic, type II {(NM_000098)
700 012	"c.110_111dupGC", "c.1239_1240delGA"}
798 - 812	CRB1:Leber congenital amaurosis 8 {(NM_201253.3) "c.1148G>A",
	"c.1576C>T", "c.1733T>A", "c.1842delT", "c.1844G>T", "c.2230C>T",
	"c.2234C>T", "c.2498G>A", "c.2555T>C", "c.2680_2684delAACCC",
	"c.3307G>A", "c.4005+1G>A", "c.4121_4130delCAACTCAGGG",
042 046	"c.424G>T", "c.455G>A"}
813 - 816	CRB2:Ventriculomegaly with cystic kidney disease {(NM_173689.7)
047 040	"c.1882C>T", "c.1928A>C", "c.2277G>A", "c.2400C>G"}
817 - 818	CRTAP:Osteogenesis imperfecta, type VII {(NM_006371) "c.976C>T"
212 212	(NM_006371.4) "c.793+1G>T"}
819 - 819	CSTA:Exfoliative ichthyosis, Autosomal Recessive, ichthyosis bullosa
	of Siemens-like {(NM_005213.4) "c.67-2A>T"}
820 - 824	CTNS:Cystinosis,CTNS-related {(NM_004937) "c.587dupA",
	"c.691C>T", "c.890G>A" (NM_004937.2) "c.1015G>A", "c.530A>C"}

825 - 825	CTSC:Haim-Munk syndrome {(NM_001814.6) "c.857A>G"}
826 - 826	CTSK:Pycnodysostosis {(NM_000396.4) "c.990A>G"}
827 - 831	CYBA:Chronic granulomatous disease, autosomal, due to deficiency of CYBA {(NM_000101) "c.160_161insC" (NM_000101.4) "c.164C>G",
	"c.171dupG", "c.70G>A", "c.71G>A"}
832 - 836	CYBB:Chronic granulomatous disease, X-linked {(NM_000397)
032 - 030	"c.1016dupC", "c.1081T>C", "c.271C>T", "c.676C>T",
	"c.90_92delCCGinsGGT"}
837 - 838	CYP11A1:Adrenal insufficiency, congenital, with 46XY sex reversal,
	partial or complete {(NM_000781.3) "c.644T>C", "c.694C>T"}
839 - 839	CYP11B2:Hypoaldosteronism, congenital, due to CMO II deficiency
	{(NM_000498.3) "c.541C>T"}
840 - 842	CYP1B1:Glaucoma 3A, primary open angle, congenital, juvenile, or
	adult onset {(NM_000104) "c.1405C>T" (NM_000104.3)
	"c.1568G>A", "c.182G>A"}
843 - 847	CYP27A1:Cerebrotendinous xanthomatosis {(NM_000784.4)
	"c.1016C>T", "c.1184G>A", "c.355delC", "c.819delT", "c.845-1G>A"}
848 - 848	CYP4F22:Congenital recessive ichthyoses (CRI) {(NM_173483)
040 040	"c.429dupG"}
849 - 849	CYP4V2:Bietti crystalline corneoretinal dystrophy {(NM_207352) "c.1123delC"}
850 - 850	C.1123delC } CYP7B1:Spastic paraplegia 5A, Autosomal Recessive {(NM_004820.5)
850 - 850	"c.1081C>T"}
851 - 851	DAG1:Muscular dystrophy-dystroglycanopathy (congenital with brain
051 - 051	and eye anomalies), type A, 9 {(NM_001165928.3) "c.743delC"}
852 - 852	DARS2:Leukoencephalopathy with brain stem and spinal cord
	involvement and lactate elevation {(NM_018122.5) "c.492+2T>C"}
853 - 854	DBT:Maple syrup urine disease, type II {(NM_001918) "c.581C>G",
	"c.939G>C"}
855 - 856	DCAF17:Woodhouse-Sakati syndome {(NM_025000) "c.580C>T"
	(NM_025000.4) "c.436delC"}
857 - 857	DCLRE1C:Severe combined immunodeficiency, Athabascan type
050 050	{(NM_001033858.2) "c.1307_1308insAGGATGCT"}
858 - 858	DDR2:Spondylometaepiphyseal dysplasia, short limb-hand type
859 - 859	{(NM_006182.4) "c.2254C>T"} DDRGK1:Spondyloepimetaphyseal dysplasia (Shohat-type)
033 - 033	{(NM_023935) "c.408+1G>A"}
860 - 860	DDX11:Warsaw breakage syndrome {(NM_030653.3) "c.1763-1G>C"}
861 - 861	DGAT1:Diarrhea 7, congenital {(NM_012079.6) "c.751+2T>C"}
862 - 863	DGUOK:Mitochondrial DNA depletion syndrome (hepatocerebral
302 300	type) {(NM_080916.3) "c.255delA", "c.271delA"}
864 - 864	DHCR24:Desmosterolosis {(NM_014762.4) "c.307C>T"}
865 - 881	DHCR7:Smith Lemli Opitz syndrome {(NM_001360) "c.1055G>A",
	"c.1342G>A" (NM_001360.2) "c.1054C>T", "c.1210C>T",
	"c.1228G>A", "c.1A>G", "c.278C>T", "c.452G>A", "c.453G>A",
	"c.506C>T", "c.724C>T", "c.725G>A", "c.755A>G", "c.906C>G", "c.964-
	1G>C", "c.964-1G>T", "c.976G>T"}
882 - 882	DHDDS:Retinitis pigmentosa 59 {(NM_024887.3) "c.124A>G"}

222 225	
883 - 886	DLD:Dihydrolipoamide Dehydrogenase Deficiency {(NM_000108.5)
007 007	"c.104dupA", "c.1123G>A", "c.1436A>T", "c.685G>T"}
887 - 887	DLL3:Spondylocostal dysostosis 1, Autosomal Recessive {(NM_016941.3) "c.395delG"}
888 - 889	DNAH11:Ciliary dyskinesia, primary, 7, with or without situs inversus
000 - 003	{(NM_001277115.2) "c.11929G>T", "c.13242_13245delAAAG"}
890 - 891	DNAH5:Ciliary dyskinesia, primary, 3, with or without situs inversus
030 - 031	(CILD3/PCD) {(NM_001369.2) "c.7502G>C", "c.8011-2A>G"}
892 - 892	DNAI1:Ciliary dyskinesia, primary, 1, with or without situs inversus
032 - 032	{(NM_012144.4) "c.1490G>A"}
893 - 894	DNAI2:Ciliary dyskinesia, primary, 9, with or without situs inversus
	{(NM_023036.6) "c.1304G>A", "c.1494+1G>A"}
895 - 895	DNAL1:Ciliary dyskinesia, primary, 16 {(NM_031427.4) "c.449A>G"}
896 - 896	DOCK8:Hyper-IgE recurrent infection syndrome, autosomal recessive
	{(NM_203447) "c.5132C>A"}
897 - 898	DOLK:Congenital disorder of glycosylation, type Im {(NM_014908.3)
	"c.1222C>G", "c.912G>T"}
899 - 900	DSG1:Erythroderma, congenital, with palmoplantar keratoderma,
	hypotrichosis, and hyper IgE {(NM_001942.4) "c.1861delG",
	"c.395C>A"}
901 - 901	DST:Epidermolysis bullosa simplex, Autosomal Recessive 2
	{(NM_183380.3) "c.14865delA"}
902 - 902	DSTYK:Spastic paraplegia, complicated {(NM_015375) "4-
	kbdeletion/20-bpinsertion"}
903 - 908	DYSF:Muscular dystrophy, limb-girdle, type 2B {(NM_003494.4)
	"c.2372C>G", "c.2779delG", "c.4741C>T",
	"c.4872_4876delGCCCGinsCCCC", "c.5057+5G>A", "c.5429G>A"}
909 - 909	ECHS1:Mitochondrial short-chain enoyl-CoA hydratase 1 deficiency
010 010	{(NM_004092) "c.476A>G"}
910 - 910	ECM1:Urbach-Wiethe disease {(NM_004425) "c.70+1G>C"}
911 - 911	EDAR:Ectodermal dysplasia 10B, hypohidrotic/hair/tooth type,
012 014	Autosomal Recessive {(NM_022336) "c.259T>C"}
912 - 914	ELP1:Dysautonomia, familial {(NM_003640.5) "c.2087G>C",
915 - 915	EOGT:Adams-Oliver syndrome 4 {(NM_001278689.2) "c.1074delA"}
916 - 919	EPG5:Vici syndrome {(NM_020964) "c.1007A>G", "c.3446G>A",
310 - 313	"c.5993C>G" (NM_020964.3) "c.5704dupT"}
920 - 920	EPM2A:Epilepsy, progressive myoclonic 2A (Lafora) {(NM_005670)
320 320	"56_kb_inclex2"}
921 - 921	ERBB3:Lethal congenital contractural syndrome 2 {(NM_001982.3)
	"c.1184-9A>G"}
922 - 922	ERCC2:Xeroderma pigmentosum, group D {(NM_000400.3)
	"c.2048G>A"}
923 - 923	ERCC5:Xeroderma pigmentosum/Cockayne {(NM_000123.3)
	"c.205C>T"}
924 - 924	ERCC6:Cockayne syndrome, type B {(NM_000124.4)
	"c.1034_1035insT"}
925 - 927	ERCC8:Cockayne syndrome, type A {(NM_000082.3) "c.37G>T",

	"c.843+1G>C", "c.966C>A"}
928 - 928	ESCO2:Roberts-SC phocomelia syndrome {(NM_001017420.3)
320 320	"c.1674-2A>G"}
929 - 932	ETFDH:Glutaric acidemia IIC {(NM_004453.4) "c.1074G>C",
323 332	"c.1084G>A", "c.1425C>A", "c.299T>A"}
933 - 933	EXOSC3:Pontocerebellar hypoplasia, type 1B {(NM_016042.4)
333 - 333	"c.571G>T"}
934 - 934	EXOSC8:Pontocerebellar hypoplasia, type 1C {(NM_181503.3)
334 - 334	"c.5C>T"}
935 - 950	EYS:Retinitis pigmentosa 25 {(NM_001142800.2) "400kb deletion in
333 - 330	6q12", "c.1211dupA", "c.3699delG", "c.3715G>T",
	"c.403_423delinsCTTTT", "c.403delA", "c.410_424del15",
	"c.4361_4362delinsAG", "c.5450G>A", "c.5928-
	37922_6078+38716del", "c.6976C>T", "c.8155_8156delCA",
	"c.8168delA", "c.8216_8217delAC", "c.8231del", "c.9286_9295del10"}
951 - 952	F7:Factor VII deficiency {(NM_000131) "c.1109G>T" (NM_000131.4)
	"c.1256C>T"}
953 - 953	FA2H:Spastic paraplegia 35, Autosomal Recessive {(NM_024306.5)
	"c.786+1G>A"}
954 - 960	FAH:Tyrosinemia, type I {(NM_000137.2) "c.1062+5G>A",
	"c.1069G>T", "c.192G>T", "c.554-1G>T", "c.707-1G>C", "c.782C>T",
	"c.786G>A"}
961 - 966	FAM161A:Retinitis pigmentosa 28 {(NM_001201543.2) "c.1003C>T",
	"c.1309A>T", "c.1321dupC", "c.1355_1356delCA", "c.1567C>T",
	"c.1786C>T"}
967 - 967	FAM20A:Amelogenesis imperfecta, type IG (enamel-renal syndrome)
	{(NM_017565.4) "c.1523delC"}
968 - 976	FANCA: Fanconi anemia, complementation group A {(NM_000135)
	"c.3382C>T" (NM_000135.4) "c.189+1G>A", "c.2172dupG",
	"c.3788_3790delTCT", "c.4168-2A>C", "c.4261-2A>C", "c.4275delT",
	"c.891_893+1delCTGG", "c.Del_exon_31-37"}
977 - 983	FANCC:Fanconi anemia, complementation group C {(NM_000136)
	"c.8_9delAA" (NM_000136.3) "c.1642C>T", "c.1661T>C", "c.37C>T",
	"c.456+4A>T", "c.553C>T", "c.67delG"}
984 - 985	FANCG:Fanconi Anemia - complementation group G {(NM_004629.1)
000 000	"c.212T>C", "c.510+3A>G"}
986 - 986	FDX1L:Mitochondrial muscle myopathy {(NM_001031734.4) "c.10A>T"}
007 000	J
987 - 989	FERMT1:Kindler syndrome {(NM_017671.4) "c.019+470del ", "c.137_140delTAGT", "c.749G>A"}
990 - 990	
990 - 990	FGB:Afibrinogenemia congenital {(NM_005141.4) "c.1400G>A"} FH:Fumarase deficiency, leiomyomatosis and renal cell cancer
771 - 771	{(NM_000143.3) "c.905-1G>A"}
992 - 994	FKBP10:Osteogenesis imperfecta, type XI {(NM_021939)
J32 - 334	"c.1271_1272delCCinsA", "c.391+4A>T" (NM_021939.3) "c.310C>T"}
995 - 995	FKRP:Muscular dystrophy-dystroglycanopathy (limb-girdle), type C, 5
- 555 - 555 	{(NM_024301.5) "c.160C>T"}
996 - 996	FKTN:Muscular dystrophy-dystroglycanopathy (congenital with brain
230 - 230	TRI William dystrophy-dystrogrycanopathy (congenital with brain

	1 1 1 1 1 1 1 1 1 1
	and eye anomalies), type A, 4 - Walker Warburg syndrome
	{(NM_001079802.1) "c.1167dupA"}
997 - 997	FLT4:Autosomal Recessive Hereditary Lymphedema {(NM_182925.5)
	"c.3704C>G"}
998 - 998	FOXRED1:Mitochondrial encephalomyopathy complex I deficiency
	{(NM_017547.4) "c.1054C>T"}
999 - 999	FRMD4A:Microcephaly intellectual disability and dysmorphism
	{(NM_018027) "c.2134_2146dup13"}
1000 - 1000	FTO:Growth retardation, developmental delay, coarse facies, and
	early death {(NM_001080432.3) "c.947G>A"}
1001 - 1002	G6PC3:Neutropenia, severe congenital 4, Autosomal Recessive
	{(NM_138387.3) "c.765_766delAG", "c.785G>A"}
1003 - 1014	G6PC:Glycogen storage disease Ia - GDS1a {(NM_000151.4)
	"c.1039C>T", "c.247C>T", "c.248G>A", "c.379_380dupTA", "c.497T>G",
	"c.508C>T", "c.562G>C", "c.648G>T", "c.724C>T", "c.79delC",
	"c.809G>T", "c.979_981delTTC"}
1015 - 1027	GAA:Pompe (Glycogen storage disease type II) {(NM_000152)
	"c.1001G>A", "c.2456G>T" (NM_000152.5) "c.1064T>C",
	"c.1082C>T", "c.1210G>A", "c.1564C>A", "c.1935C>A", "c.1942G>A",
	"c.2560C>T", "c.340_341insT", "c.670C>T", "c.896T>C", "c.896T>G"}
1028 - 1030	GALC:Krabbe disease {(NM 000153.4) "c.1630G>A", "c.1748A>C",
	"c.1796T>G"}
1031 - 1032	GALNT3:Tumoral calcinosis, hyperphosphatemic, familial
	{(NM_004482.4) "c.1524+1G>A", "c.1524+5G>A"}
1033 - 1042	GALT:Galactosemia {(NM_000155.3) "5.5-KB_DEL", "c.152G>A",
1033 1042	"c.253-2A>G", "c.404C>T", "c.413C>T", "c.512T>C", "c.563A>G",
	"c.584T>C", "c.626A>G", "c.855G>T"}
1043 - 1044	GAN:Giant axonal neuropathy 1 {(NM_022041) "c.103G>T"
	(NM_022041.3) "c.973G>A"}
1045 - 1045	GATC:Hypertophic Cardiomyopathy {(NM_176818) "c.233T>G"}
1046 - 1046	GATM:Cerebral creatine deficiency syndrome 3 {(NM_001482.3)
1040 - 1040	"c.1111dupA"}
1047 - 1058	GBA:Gaucher disease, , type I {(NM_001005741.3) "c.115+1G>A",
1047 - 1036	"c.1226A>G", "c.1294T>A", "c.1297G>T", "c.1342G>C", "c.1448T>C",
	"c.1504C>T", "c.1505G>A", "c.1604G>A", "c.259C>T", "c.703T>C",
	"c.84dupG"}
1059 - 1070	GCDH:Glutaricaciduria type I {(NM_000159.4) "c.1168G>C",
1039 - 1070	"c.1173delG", "c.1204C>T", "c.1205G>A", "c.1247C>T", "c.1262C>T",
	"c.1306G>T", "c.301G>A", "c.505+1G>A", "c.848T>C", "c.877G>A",
	"c.914C>T"}
1071 1072	
1071 - 1072	GH1:Growth hormone deficiency, isolated, type IA {(NM_000515.5) "c.456+5G>C", "c.67G>T"}
4072 4072	· ,
1073 - 1078	GHR:Laron dwarfism {(NM_000163.5) "c.11G>A", "c.594A>G",
4070 4070	"c.62G>A", "c.703C>T", "c.744delT", "del5,6ex"}
1079 - 1079	GHRHR:Growth hormone deficiency, isolated, type IB
1000 1000	{(NM_000823.4) "c.1069C>T"}
1080 - 1080	GIPC3:Deafness, autosomal recessive 15 {(NM_133261) "c.937T>C"}
1081 - 1095	GJB2:Deafness, autosomal recessive 1A {(NM_004004.6) "c.109G>A",

	"c.167delT", "c.229T>C", "c.230G>A", "c.235delC", "c.250G>A",
	"c.269T>C", "c.358_360delGAG", "c.35delG", "c.370C>T",
	"c.51_62delCACCAGCATTGGinsA", "c.551G>C", "c.614T>C", "c.71G>A", "c.94C>T"}
1096 - 1096	GJB6:Deafness, Autosomal Recessive 1B {(NM_006783.4) "309_kb"}
1097 - 1102	GLB1:GM1-gangliosidosis, type I {(NM_000404.4) "c.1038G>C",
	"c.485delT", "c.602G>A", "c.824A>G", "c.827A>C", "c.914+4A>G"}
1103 - 1106	GLDC:Glycine encephalopathy and non-ketoic hyperglycinemia, GLDC-
	related {(NM_000170.2) "c.2405C>T", "c.2607C>A", "c.2T>C",
	"c.985C>A"}
1107 - 1107	GLRA1:Hyperekplexia, hereditary 1, autosomal dominant or recessive
	{(NM_001146040.1) "c.298C>T"}
1108 - 1108	GMPPA:Alacrima, achalasia, and mental retardation syndrome
	{(NM_013335.3) "c.1000A>C"}
1109 - 1110	GMPPB:Muscular dystrophy-dystroglycanopathy {(NM_013334.3)
	"c.656T>C", "c.860G>A"}
1111 - 1111	GNE:Hereditary inclusion body myopathy (HIBM) {(NM_005476.6)
	"c.2135T>C"}
1112 - 1118	GNPTAB:Mucolipidosis III alpha/beta {(NM_024312.5) "c.118-2A>G",
	"c.2314_2315insA", "c.2918dupT", "c.3434+1G>A", "c.3434+715G>A",
	"c.3503_3504delTC", "c.3613C>T"}
1119 - 1119	GNPTG:Mucolipidosis III gamma {(NM_032520.5) "c.499dupC"}
1120 - 1121	GPC6:Omodysplasia 1 {(NM_005708)
	"g.93997007_94063501del66495insATAAATCACTTAGAGATGT",
	"g.94252984_94352299del99316insCTA"}
1122 - 1122	GPSM2:Chudley-McCullough syndrome {(NM_013296.5) "c.379C>T"}
1123 - 1123	GRHPR:Hyperoxaluria, primary, type II {(NM_012203.2) "c.975A>G"}
1124 - 1131	GUCY2D:Leber congenital amaurosis 1, Cone-rod dystrophy 6
	{(NM_000180.3) "c.1992T>G", "c.2129C>T", "c.2513G>A",
	"c.2618C>G", "c.389delC", "c.529C>T", "c.620delC", "c.693delG"}
1132 - 1132	HACD1:Congenital myopathy {(NM_014241.4) "c.744C>A"}
1133 - 1133	HADHA:Long-Chain hydroxylacyl-CoA dehydrogenase deficiency
	(LCHAD) {(NM_000182.5) "c.1528G>C"}
1134 - 1134	HAX1:Severe congenital neutropenia type 3 (SCN3), a.k.a. Kostmann
	disease {(NM_006118) "c.125dupG"}
1135 - 1161	HBB:Hemoglobipathies (Including sickle-cell anemia and beta
	thalassemia, Hb C, D, E, O) {(NM_000518) "c138C>A", "c50-
	101C>T", "c78A>C", "c80T>A", "c.112delT", "c.114G>A",
	"c.118C>T", "c.135delC", "c.19G>A", "c.315+1G>A", "c.364G>A",
	"c.364G>C", "c.79G>A", "c.82G>T", "c.92+5G>C", "c.92+6T>C",
	"c.92G>C", "c.93-22_95del25" (NM_000518.5) "1.78_Mb",
	"c.17_18delCT", "c.20A>T", "c.25_26delAA", "c.27dupG", "c.316-
	106C>G", "c.47G>A", "c.92+1G>A", "c.93-21G>A"}
1162 - 1184	HEXA:Tay-Sachs disease {(NM_000520) "c.1176G>A", "c.1528C>T"
	(NM_000520.5) "c.1073+1G>A", "c.1274_1277dupTATC",
	"c.1351C>G", "c.1421+1G>C", "c.1444G>A", "c.316C>T",
	"c.459+2dupT", "c.496delC", "c.509G>A", "c.532C>T", "c.533G>A",
	"c.533G>T", "c.540C>G", "c.571-2A>G", "c.749G>A", "c.749G>T",

	700, A 00f, 10, A 00f, A 02fm, C
	"c.78G>A", "c.805+1G>A", "c.805G>A", "c.835T>C", "c.910_912delTTC"}
1185 - 1185	HEXB:Sandhoff disease, infantile, juvenile, and adult forms
1103 - 1103	{(NM_000521) "c.1082+5G>A"}
1186 - 1186	HGD:Alkaptonuria {(NM_000187) "c.16-272_87+305del"}
1187 - 1187	HGSNAT:Retinitis pigmentosa 73 {(NM_152419.3) "c.370A>T"}
1188 - 1188	HIKESHI:Leukodystrophy, early onset spastic paraparesis, acquired
1100 1100	microcephaly, optic atrophy and risk of early death {(NM_016401.4)
	"c.160G>C"}
1189 - 1191	HMGCL:HMG-CoA lyase deficiency {(NM_000191.3) "c.122G>A",
	"c.125A>G", "c.521G>A"}
1192 - 1192	HOGA1:Hyperoxaluria, primary, type III {(NM_138413)
	**"c.944_946delAGG"}
1193 - 1195	HPD:Thyrosinemia type III {(NM_002150.3) "c.325-1G>A", "c.415-
1105 1105	1G>A", "c.481G>C"}
1196 - 1196 1197 - 1200	HPS1:Hermansky-Pudlak syndrome 1 {(NM_000195.5) "c.972delC"} HPS3:Hermansky-Pudlak syndrome 3 {(NM_032383.5) "c
1197 - 1200	2993_217+690del3900", "c.1163+1G>A", "c.1691+2T>G", "c.2482-
	2A>G"}
1201 - 1201	HPS6:Hermansky-Pudlak syndrome 6 {(NM_024747.5) "c.1065dupG"}
1202 - 1202	HSPD1:Leukodystrophy, hypomyelinating, 4, HLD4 (HSP60)
	{(NM_199440.1) "c.86A>G"}
1203 - 1203	IBA57:Spastic paraplegia 74, Autosomal Recessive
	{(NM_001010867.4) "c.678A>G"}
1204 - 1207	IDUA:Mucopolysaccharidosis Type IH - Hurler syndrome
	{(NM_000203.5) "c.1096A>C", "c.192C>A", "c.208C>T", "c.928C>T"}
1208 - 1209	IGHMBP2:Neuronopathy, distal hereditary motor, type VI
1210 - 1210	{(NM_002180.2) "c.114delA", "c.707T>G"} IL10RA:Inflammatory bowel disease 28, early onset, autosomal
1210 - 1210	recessive {(NM_001558) "c.537G>A"}
1211 - 1214	INSR:Leprechaunism, Donohue syndrome {(NM_000208) "c.2683-
	542_2842+544del" (NM_000208.4) "c.167T>C", "c.3079C>T",
	"c.857G>A"}
1215 - 1215	INVS:Nephronophthisis 2, infantile {(NM_014425.5) "c.2719C>T"}
1216 - 1216	ISPD:Muscular dystrophy-dystroglycanopathy (congenital with brain
	and eye anomalies), {(NM_001101426.4) "c.165dupG"}
1217 - 1220	ITGA2B:Glanzmann thrombasthenia, ITGA2B-related {(NM_000419)
	"c.2374delG" (NM_000419.4) "c.1947-1G>A", "c.818G>A", "c.97A>G"}
1221 - 1223	ITGB3:Glanzmann thrombasthenia, ITGB3-related {(NM 000212)
1221 - 1223	"c.1616 1617delTT" (NM 000212.2) "11.2kbincl.ex.10-partex.13",
	"c.428T>G"}
1224 - 1224	ITGB4:Epidermolysis bullosa, junctional, with pyloric atresia - Carmi
	syndrome {(NM_000213.5) "c.3280_3793+176del2279 "}
1225 - 1225	ITK:Lymphoproliferative syndrome {(NM_005546) "c.1764C>G"}
1226 - 1229	IVD:Isovaleric academia {(NM_002225.4) "c.148C>T", "c.286+2T>C",
	"c.456+2T>C", "c.932C>T"}

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1230 - 1230	JAK3:SCID, autosomal recessive, T-negative/B-positive type {(NM_000215) "c.2680+89G>A"}
1231 - 1231	KCNJ10:SESAME syndrome {(NM_002241.5) "c.524G>A"}
1232 - 1232	KIAA1279:Goldberg-Shprintzen megacolon syndrome {(NM_015634)
1232 1232	"c.1516dupA"}
1233 - 1233	KIF1C:Spastic ataxia 2, Autosomal Recessive {(NM_006612)
	"c.2191C>T"}
1234 - 1234	KIZ:Retinitis pigmentosa 69 {(NM_018474) "c.226C>T"}
1235 - 1235	KLHL40:Nemaline myopathy 8, Autosomal Recessive {(NM_152393.4) "c.581T>A"}
1236 - 1236	KREMEN1:Ectodermal dysplasia {(NM_032045) "c.626T>C"}
1237 - 1238	KRT14:Epidermolysis bullosa simplex {(NM_000526) "c.400C>T", "c.915G>A"}
1239 - 1240	KY:Myopathy, myofibrillar, 7 {(NM_178554) "c.405C>A",
	"c.51_52insTATCGACATGTGCTGTATCTATCGACAT"}
1241 - 1246	LAMA2:Muscular dystrophy, congenital, due to partial LAMA2
	deficiency {(NM_000426) "c.4609_4631del" (NM_000426.3)
1247 - 1250	"c.3718C>T", "c.5260delG", "c.828C>G", "c.8665G>A", "c.8689C>T"}
	LAMA3:Laryngoonychocutaneous Syndrome {(NM_000227.4)
1251 - 1261	LAMB3:Epidermolysis bullosa, junctional, non-Herlitz type
	{(NM_000228) "c.129insA" (NM_000228.3) "c.124C>T",
	"c.1295dupA", "c.1903C>T", "c.1978C>T", "c.2528delA", "c.2914C>T", "c.3024delT", "c.3247C>T", "c.430C>T", "c.727C>T"}
1262 - 1263	LAMC2:Epidermolysis bullosa, junctional, Herlitz type
1202 1203	{(NM_018891.2) "c.1756C>T", "c.368_373delinsACCAC"}
1264 - 1268	LCA5:Leber congenital amaurosis 5 {(NM_181714.3)
	"c.1062_1068delCGAAAAC", "c.1714C>T", "c.238C>T", "c.835C>T",
	"c.94delT"}
1269 - 1270	LIFR:Stuve-Wiedemann syndrome/Schwartz-Jampel type 2
	syndrome-LIFR related {(NM_002310.5) "c.1601-1G>A",
1271 - 1272	LIPA:Wolman disease {(NM_001127605.2) "c.260G>T", "c.398delC"}
1273 - 1273	LONP1:CODAS syndrome {(NM_004793.4) "c.2009C>T"}
1274 - 1275	LOXHD1:Deafness, Autosomal Recessive 77 {(NM_144612)
	"c.5894dupG" (NM_144612.6) "c.4714C>T"}
1276 - 1277	LRBA:Immunodeficiency, common variable, 8, with autoimmunity
	{(NM_001199282) "c.8139_8142dupCATG" (NM_001199282.2)
	"c.7937T>G"}
1278 - 1279	MAK:Retinitis pigmentosa 62 {(NM_001242957.2) "c.497G>A"
1200 1200	(NM_005906) "c.394_395insCTTC"}
1280 - 1280	MAN1B1:Mental retardation, Autosomal Recessive 15 {(NM_016219.5) "c.1863G>A"}
1281 - 1281	MATN3:Spondyloepimetaphyseal dysplasia {(NM_002381.5)
	"c.910T>A"}
1282 - 1282	MCIDAS:Mucociliary clearance disorder {(NM_001190787.2)
	"c.1142G>A"}
1283 - 1287	MCOLN1:Mucolipidosis type IV - ML4 {(NM_020533) "c.1135-1G>C"

	(NIX 000500 0) 4045 500 16400 40050 50 406 04 0
	(NM_020533.3) "c1015_788del6433", "c.1207C>T", "c.406-2A>G", "c.964C>T"}
1288 - 1289	MECR:Dystonia, childhood-onset, with optic atrophy and basal ganglia abnormalities {(NM_016011) "c.695G>A", "c.830+2dupT"}
1290 - 1290	MED17:Microcephaly, postnatal progressive, with seizures and brain atrophy ((ICCA) {(NM_004268.5) "c.1112T>C"}
1291 - 1291	MED25:Basel-Vanagaite-Smirin-Yosef syndrome {(NM_030973.3) "c.116A>G"}
1292 - 1292	MEGF10:Myopathy, areflexia, respiratory distress, and dysphagia, early-onset {(NM_001256545.2) "c.1325delC"}
1293 - 1293	MERTK:Retinitis pigmentosa 38 {(NM_006343) "c.2164C>T"}
1294 - 1295	MFSD8:Ceroid lipofuscinosis, neuronal, 7 {(NM_152778) "c.103C>T" (NM_152778.2) "c.472G>A"}
1296 - 1297	MKS1:Meckel syndrome 1 {(NM_017777.3) "c.1048C>T", "c.472C>T"}
1298 - 1300	MLC1:Megalencephalic leukoencephalopathy with subcortical cysts {(NM_015166.3) "c.176G>A", "c.274C>T", "c.278C>T"}
1301 - 1301	MLPH:Griscelli syndrome, type 3 {(NM_024101.7) "c.103C>T"}
1302 - 1302	MMACHC:Methylmalonic aciduria and homocystinuria, cblC type {(NM_015506.3) "c.271dupA"}
1303 - 1305	MOCS1:Molybdenum cofactor deficiency A {(NM_001075098.3) "c.1510C>T", "c.722delT", "c.971G>A"}
1306 - 1307	MOCS2:Molybdenum cofactor deficiency Type B {(NM_004531.5) "c.226G>A", "c.377+1G>A"}
1308 - 1311	MPDU1:Congenital disorder of glycosylation, type If {(NM_004870)
1312 - 1317	MPL:Thrombocytopenia, congenital amegakaryocytic {(NM_005373) "c.212+5G>A", "c.76C>T" (NM_005373.2) "c.1031T>A", "c.127C>T", "c.460T>C", "c.79+2T>A"}
1318 - 1318	MPV17:Mitochondrial DNA depletion syndrome 6 (hepatocerebral type) {(NM_002437.5) "c.278A>C"}
1319 - 1319	MRE11A:Ataxia Telangiectasia like disorder {(NM_005591.3) "c.290A>G"}
1320 - 1322	MTHFR:Homocystinuria due to MTHFR deficiency {(NM_005957) "c.1072C>T" (NM_005957.4) "c.16delA", "c.474A>T"}
1323 - 1326	MTTP:Abetalipoproteinemia ABL {(NM_000253.3) "c.2212delT", "c.2593G>T", "c.307A>T", "c.62-2A>G"}
1327 - 1328	MUT:Methylmalonic acidemia, mut(0) type {(NM_000255) "c.1240G>T" (NM_000255.4) "c.655A>T"}
1329 - 1329	MVK:Hyper-IgD syndrome {(NM_000431.4) "c.1129G>A"}
1330 - 1332	MYBPC1:Lethal congenital contracture syndrome 4 {(NM_002465.4) "c.556G>A", "c.688G>A", "c.952C>T"}
1333 - 1334	MYH2:Proximal myopathy and ophthalmoplegia {(NM_017534.6) "c.2400delG", "c.706G>A"}
1335 - 1341	MY015A:Deafness, Autosomal Recessive 3 {(NM_016239) "c.1223C>T", "c.9861C>T" (NM_016239.4) "c.373_374delCG", "c.4240G>A", "c.7207G>T", "c.8183G>A", "c.8467G>A"}
1342 - 1361	MYO7A:Usher syndrome, type 1B {(NM_000260) "c.3262C>T" (NM_000260.4) "c.1190C>A", "c.1996C>T", "c.2065delC",

	"c.2187+1G>A", "c.2307delC", "c.2476G>A", "c.2777T>A", "c.29T>C", "c.3892G>A", "c.4153-2A>G", "c.470+1G>A", "c.5068dupC",
	"c.5581C>T", "c.5660C>T", "c.6196delC", "c.620A>G", "c.640G>A",
	"c.6487G>A", "c.700C>T"}
1362 - 1362	NAGLU:Mucopolysaccharidosis type IIIB (Sanfilippo B)
	{(NM_000263.4) "c.2021G>A"}
1363 - 1363	NARS2:Combined oxidative phosphorylation deficiency 24
	(COXPD24) {(NM_024678) "c.500A>G"}
1364 - 1365	NBEAL2:Gray platelet syndrome {(NM_015175.2) "c.2701C>T", "c.5413dupG"}
1366 - 1368	NCF1:Chronic granulomatous disease due to deficiency of NCF-1
	{(NM_000265) "c.75_76delGT" (NM_000265.6) "c.153+1G>A",
1369 - 1372	*"c.579G>A"} NCF2:Chronic granulomatous disease due to deficiency of NCF-2
1309 - 1372	{(NM 000433) "exon9+10insertionafterexon10" (NM 000433.3)
	"c.1171_1175delAAGCT", "c.196C>T", "c.304C>T"}
1373 - 1373	NDUFA11:Mitochondrial complex I deficiency - NDUFA11 gene
	{(NM_001193375.1) "c.97+5G>A"}
1374 - 1374	NDUFAF5:Mitochondrial complex I deficiency - NDUFAF5 gene
	{(NM_024120.5) "c.749G>T"}
1375 - 1375	NDUFS2:Mitochondrial complex I deficiency-NDUFS2 gene
1376 - 1376	{(NM_004550.4) "c.1237T>C"} NDUFS4:Leigh syndrome {(NM_002495.4) "c.462delA"}
1377 - 1377	NDUFS6:Mitochondrial complex I deficiency - NDUFS6 gene
13// - 13//	{(NM_004553.4) "c.344G>A"}
1378 - 1381	NEB:Nemaline myopathy 2 {(NM_001271208.2) "c.17118+1G>A",
	"c.18808C>T", "c.9619-2A>G" (NM_004543.4)
	"c.7431+1917_7536+372del"}
1382 - 1382	NECTIN1:Cleft lip/palate ectodermal dysplasia, CLPED1 (Zlotogora- Ogur syndrome) {(NM_203285) "c.556delG"}
1383 - 1383	NGLY1:Congenital disorder of deglycosylation {(NM_018297.4)
	"c.1294G>T"}
1384 - 1384	NNT:Glucocorticoid deficiency 4 {(NM_182977.3) "c.598G>A"}
1385 - 1403	NPC1:Niemann-Pick disease type C1 {(NM_000271) "c.3742_3753del" (NM_000271.5) "c.1211G>A", "c.1241_1242delTC",
	(NM_000271.5)
	"c.2279_2281delTCT", "c.2780C>T", "c.2972_2973delAG",
	"c.2974G>A", "c.2974G>C", "c.2974G>T", "c.3007C>T",
	"c.3347_3348delTC", "c.3467A>G", "c.3557G>A", "c.3614C>A",
	"c.3637T>G", "c.3673T>G"}
1404 - 1404	NPHP1:Joubert syndrome {"del exons 2-7"}
1405 - 1413	NPHS1:Nephrotic syndrome type 1 {(NM_004646.3) "c.1138C>T",
	"c.121_122delCT", "c.1707C>G", "c.2104G>A", "c.2160dupC",
1414 - 1415	"c.3325C>T", "c.3478C>T", "c.514_516delACC", "c.532C>T"} NPHS2:Nephrotic syndrome {(NM_014625) "c.388G>A"
1414 - 1412	(NM_014625.3) "c.412C>T"}
1416 - 1417	NRL:Retinitis pigmentosa 27 {(NM_006177)
	"c.444_445insGCTGCGGG", "c.91C>T"}

1418 - 1421	NTRK1:Insensitivity to pain, congenital, with anhidrosis (CIPA) {(NM_002529.3) "c.1250C>T", "c.1860_1861insT", "c.207_208delTG", "c.2084C>T"}
1422 - 1422	NUP62:Striatonigral degeneration, Infantile Bilateral Striatal Necrosis (IBSN) {(NM_016553.4) "c.1172A>C"}
1423 - 1423	OAT:Gyrate atrophy of choroid and retina with or without ornithinemia {(NM_000274) "c.159delC"}
1424 - 1426	OCA2:Albinism, oculocutaneous, type II {(NM_000275) "c.79G>A" (NM_000275.3) "c.1320G>C", "c.1327G>A"}
1427 - 1427	OPA3:3-methylglutaconic aciduria, type III - Costeff {(NM_025136.3) "c.143-1G>C"}
1428 - 1430	OTC:Ornithine transcarbamylase deficiency {(NM_000531.6) "c.717+1G>T", "c.829C>T", "c.958C>T"}
1431 - 1432	OTOA:Deafness, Autosomal Recessive 22 {(NM_144672) "c.1025A>T", "c.2359G>T"}
1433 - 1434	OTOF:Deafness, Autosomal Recessive 9 {(NM_194248) "c.5332G>T" (NM_194248.2) "c.2866+1G>A"}
1435 - 1435	P3H2:Myopia, high, with cataract and vitreoretinal degeneration {(NM_018192) "c.1523G>T"}
1436 - 1475	PAH:Phenylketonuria {(NM_000277) "c.1089delG", "c.1139C>T", "c.116T>C", "c.1184C>G", "c.1243G>A", "c.168+1G>A", "c.169-4951del6604ins8 6.7kb_del", "c.169_171delGAG", "c.283A>T", "c.311C>A", "c.350delC", "c.442-5C>G", "c.48dupT", "c.506G>A", "c.526C>T", "c.592_613del22", "c.632delC", "c.838G>A", "c.842+5G>A", "c.967_969delACA", "c.969+1G>A" (NM_000277.3) "c.1045T>C", "c.1066-11G>A", "c.1208C>T", "c.1222C>T", "c.1315+1G>A", "c.143T>C", "c.165T>G", "c.165delT", "c.441+5G>T", "c.473G>A", "c.689T>C", "c.722G>A", "c.727C>T", "c.754C>T", "c.782G>A", "c.782G>C", "c.842C>T", "c.889C>T", "c.898G>T"}
1476 - 1476	PARK2:Parkinson disease, early onset {(NM_004562) "c.101delA"}
1477 - 1477	PAX7:Myopathy, congenital, progressive, with scoliosis {(NM_001135254) "c.1403-2A>G"}
1478 - 1478	PCCA:Propionic acidemia, PCCA-related {(NM_000282.4) "c.923dupT"}
1479 - 1479	PCCB:Propionic acidemia, PCCB-related {(NM_000532.5) "c.1173dupT"}
1480 - 1481	PCDH12:Microcephaly, seizures, spasticity, and brain calcification (MISSBC) {(NM_016580) "c.2515C>T", "c.995delT"}
1482 - 1482	PCDH15:Usher syndrome, type 1F {(NM_033056.3) "c.733C>T"}
1483 - 1483	PCK1:Phosphoenolpyruvate carboxykinase-1, cytosolic, deficiency {(NM_002591.4) "c.134T>C"}
1484 - 1485	PCNT:Microcephalic osteodysplastic primordial dwarfism, type II {(NM_006031) "c.2984_2994delCAGACTTTGAG" (NM_006031.5) "c.3465-1G>A"}
1486 - 1490	PDE6A:Retinitis pigmentosa 43 {(NM_000440) "c.1957C>T", "c.1960C>T", "c.2081_2085delAACAG", "c.409delGinsCT", "c.769C>T"}
1491 - 1491	PDE6B:Retinitis pigmentosa-40 {(NM_001145291) "c.1417delC"}
1492 - 1492	PDE6G:Retinitis pigmentosa 57 {(NM_002602.4) "c.187+1G>T"}

1493 - 1495	PEPD:Prolidase deficiency {(NM_000285.4) "c.1103T>G", "c.605C>T", "c.634G>C"}
1496 - 1498	PEX1:Peroxisome biogenesis disorder 1A (Zellweger) {(NM_000466) "c.2916delA" (NM_000466.3) "c.2097dupT", "c.2528G>A"}
1499 - 1501	PEX2:Peroxisome biogenesis disorder 5A (Zellweger) {(NM_001079867.1) "c.355C>T", "c.550delT", "c.669G>A"}
1502 - 1506	PEX6:Peroxisome biogenesis disorder 4B (Zellweger syndrome) {(NM_000287.4) "c.1715C>T", "c.1944delC", "c.1947delG", "c.2094+2T>C", "c.2534T>C"}
1507 - 1507	PEX7:Rhizomelic chondrodysplasia punctata type 1 {(NM_000288.4) "c.283T>G"}
1508 - 1508	PGAP3:Hyperphosphatasia with mental retardation syndrome 4 {(NM_033419.5) "c.845A>G"}
1509 - 1509	PGM1:Congenital disorder of glycosylation, type It {(NM_002633) "c.112A>T"}
1510 - 1510	PHGDH:Phosphoglycerate dehydrogenase deficiency {(NM_006623.3) "c.1468G>A"}
1511 - 1511	PHKG2:Glycogen storage disease IXc {(NM_000294.3) "c.71A>G"}
1512 - 1512	PHYH:Refsum disease {(NM_001037537.1) "c.523C>T"}
1513 - 1514	PIGN:Multiple congenital anomalies-hypotonia-seizures syndrome 1 {(NM_012327.5) "c.2126G>A", "c.755A>T"}
1515 - 1516	PIGT:Multiple congenital anomalies-hypotonia-seizures syndrome 3 {(NM_015937.6) "c.1564T>G", "c.761delG"}
1517 - 1517	PIP5K1C:Lethal congenital contractural syndrome 3 {(NM_012398.2) "c.757G>A"}
1518 - 1518	PJVK:Deafness, Autosomal Recessive 59 {(NM_001042702.4) "c.406C>T"}
1519 - 1529	PKHD1:Polycystic kidney & hepatic disease, PKHD1-related {(NM_138694) "c.10444C>T", "c.1486C>T", "c.5895dupA", "c.8870T>C", "c.9689delA" (NM_138694.4) "c.107C>T", "c.1350delC", "c.2279G>A", "c.3761_3762delCCinsG", "c.6122-12G>A", "c.824C>T"}
1530 - 1534	PLA2G6:Infantile neuroaxonal dystrophy 1 (INAD) {(NM_003560) "c.1594A>T" (NM_003560.4) "c.1040G>C", "c.2070_2072delTGT", "c.2251G>A", "c.668C>A"}
1535 - 1535	PLAA:Neurodevelopmental disorder with progressive microcephaly, spasticity, and brain anomalies {(NM_001031689.3) "c.2254C>T"}
1536 - 1536	PLEKHG2:Leukodystrophy and acquired microcephaly with or without dystonia {(NM_022835.3) "c.610C>T"}
1537 - 1540	PMM2:Congenital disorder of glycosylation Ia {(NM_000303) "c.338C>T", "c.357C>A", "c.422G>A", "c.691G>A"}
1541 - 1541	POC1A:Short stature, onychodysplasia, facial dysmorphism, and hypotrichosis {(NM_015426.5) "c.512T>C"}
1542 - 1542	POMGNT2:Muscular dystrophy-dystroglycanopathy (congenital with brain and eye anomalies, type A, 8) {(NM_032806.6) "c.1232_1233delAG"}
1543 - 1544	POMT1:Walker-Warburg Syndrome, type A, 1 {(NM_007171) "c.2167dupG", "c.428-1G>C"}

1545 - 1545	POMT2:Walker-Warburg Syndrome, type A, 2 {(NM_013382) "c.924-
4546 4546	2A>C"}
1546 - 1546	POR:Antley-Bixler syndrome with genital anomalies and disordered steroidogenesis {(NM_000941.3) "c.1615G>A"}
1547 - 1547	PPIB:Osteogenesis imperfecta, type IX {(NM_000942.4)
1547 - 1547	"c.563_566delACAG"}
1548 - 1548	PPP1R13L:Cardio-Cutaneous Syndrome DCM {(NM_006663.4)
1348 - 1348	"c.2241C>G"}
1549 - 1549	PPT1:Ceroid lipofuscinosis, neuronal, 1 {(NM_000310.3)
1545 1545	"c.169dupA"}
1550 - 1550	PRCD:Retinitis pigmentosa 36 {(NM_001077620) "c.64C>T"}
1551 - 1551	PRICKLE1:Epilepsy, progressive myoclonic 1B {(NM_153026.3)
	"c.311G>A"}
1552 - 1552	PSMB8:Autoinflammation, lipodystrophy, and dermatosis syndrome
	{(NM_148919.4) "c.405C>A"}
1553 - 1553	PTPN23:Developmental delay, cognitive impairment, and atopic
	atrophy {(NM_015466) "c.3886_3888del"}
1554 - 1554	PUS1:Mitochondrial myopathy and sideroblastic anemia 1
	{(NM_001002020.3) "c.346C>T"}
1555 - 1555	RAB27A:Griscelli syndrome, type 2 {(NM_004580)
	"c.148_149delinsC"}
1556 - 1556	RAB28:Cone-rod dystrophy 18 {(NM_001017979) "c.409C>T"}
1557 - 1558	RAG1:Severe combined immudeficiency, B cell-negative, RAG1-
	related {(NM_000448.2) "c.1361T>A", "c.1410_1413delCTTG"}
1559 - 1563	RAG2:Severe combined immudeficiency, B cell-negative, RAG2-
	related {(NM_000536.3) "c.1438G>T", "c.193G>T", "c.379A>T",
1564 - 1567	"c.470G>T", "c.685C>T"} RAPSN:Myasthenic syndrome, congenital, associated with
1504 - 1507	acetylcholine receptor deficiency {(NM_005055.5) "c210A>G", "c
	27C>G", "c.264C>A", "c.672_673insACT"}
1568 - 1568	RAPSN:Severe combined immudeficiency, B cell-negative, RAG2-
	related {(NM_005055) "c.648T>A"}
1569 - 1569	RARS2:Pontocerebellar hypoplasia, type 6 {(NM_020320.5)
	"c.110+5A>G"}
1570 - 1577	RDH12:Leber congenital amaurosis 13 {(NM_152443.3) "c.146C>T",
	"c.164C>T", "c.295C>A", "c.377C>T", "c.481C>T", "c.658+1G>A",
	"c.716G>A", "c.740T>C"}
1578 - 1579	RECQL2:Werner syndrome {(NM_000553.5) "c.1105C>T",
	"c.2665C>T"}
1580 - 1580	RFX5:Bare lymphocyte syndrome, type II (SCID) {(NM_000449)
1501 1501	"c.715C>T"}
1581 - 1581	RIN2:Macrocephaly, alopecia, cutis laxa, and scoliosis {(NM_018993.3) "c.1731delC"}
1582 - 1582	RNASEH2B:Aicardi-Goutieres syndrome 2 {(NM_024570.3)
1302 1302	"c.529G>A"}
1583 - 1583	ROGDI:Kohlschutter-Tonz syndrome {(NM_024589.2) "c.469C>T"}
1584 - 1585	RP1:Retinitis pigmentosa 1 {(NM_006269) "c.688G>T"
	(NM_006269.2) "c.4941dupT"}
	1 \(- \)

1500 1501	DDECET 1
1586 - 1591	RPE65:Leber congenital amaurosis 2 {(NM_000329.3) "c.1301C>G",
4502 4505	"c.227A>C", "c.361dupT", "c.722A>T", "c.886dupA", "c.95-2A>T"}
1592 - 1595	RPGRIP1:Cone-rod dystrophy 13 {(NM_020366) "c.1615_1624delGAACTGGAGG", "c.2935C>T", "c.2974delA",
	"c.3663_3666delAGAA"}
1596 - 1596	RPGRIP1L:Meckel syndrome 5 {(NM_015272.5) "c.118C>T"}
1597 - 1597	RRM2B:Mitochondrial DNA depletion syndrome 8 {(NM_015713.5)
1597 - 1597	"c.215C>G"}
1598 - 1598	RSPH9:Ciliary dyskinesia, primary, 12 {(NM_152732.5) "c.804_806delGAA"}
1599 - 1603	RTEL1:Dyskeratosis congenita {(NM_001283009.1) "c.1476G>T", "c.2848C>T", "c.2869C>T", "c.2920C>T", "c.3791G>A"}
1604 - 1606	RYR1:Minicore myopathy with external ophthalmoplegia {(NM_000540) "c.1366G>A", "c.9047A>G" (NM_000540.2) "c.9623C>T"}
1607 - 1608	SAMD9:Tumoral calcinosis, familial, normophosphatemic {(NM_017654.4) "c.1030C>T", "c.4483A>G"}
1609 - 1613	SAMHD1:Aicardi Goutieres syndrome {(NM_015474) "c.359_370delATCCTATCCATG" (NM_015474.3) "9.1-KB_DEL", "c.1106T>C", "c.649_650insG", "c.676C>G"}
1614 - 1614	SARS2:Hyperuricemia, pulmonary hypertension, renal failure, and alkalosis {(NM_017827.3) "c.1169A>G"}
1615 - 1615	SCAPER:Retinitis pigmentosa with intellectual disability {(NM_020843) "c.2806delC"}
1616 - 1617	SCN9A:Insensitivity to pain, congenital, with anhidrosis (CIPA) {(NM_002977.3) "c.1124delG", "c.2687G>A"}
1618 - 1618	SCNN1A:Pseudohypoaldosteronism type I - SCNN1A gene {(NM_001038) "c.1522C>T"}
1619 - 1620	SCNN1B:Pseudohypoaldosteronism type I - SCNN1B gene {(NM_000336) "c.648dupA", "c.915delC"}
1621 - 1622	SDHA:Cardiomyopathy, dilated , 1GG neonatal isolated {(NM_004168)
1623 - 1624	SEC23B:Dyserythropoietic anemia, congenital, type II {(NM_006363.6) "c.2129C>T", "c.325G>A"}
1625 - 1629	SERAC1:3-methylglutaconic aciduria with deafness, encephalopathy, and Leigh-like syndrome {(NM_032861) "c.1102C>T", "c.1339C>T" (NM_032861.4) "c.1018delT", "c.128+4A>G", "c.698_699delinsAGTATA"}
1630 - 1630	SGCG:Muscular dystrophy, limb-girdle, type 2C {(NM_000231.2) "c.525delT"}
1631 - 1639	SGSH:Mucopolysaccharidisis type IIIA (Sanfilippo A) {(NM_000199) "c.1231C>T", "c.267C>A", "c.697C>T" (NM_000199.5) "c.1093C>T", "c.1298G>A", "c.332T>C", "c.416C>T", "c.544C>T", "c.812C>T"}
1640 - 1640	SLC12A3:Bartter Syndrome, Gitelman Variant {(NM_000339.3) "c.1313G>A"}
1641 - 1641	SLC17A5:Sialic acid storage disorder, infantile (ISSD) {(NM_012434.5) "c.983G>A"}
1642 - 1642	SLC18A3:Myasthenia gravis, congenital {(NM_003055) "c.1078G>C"}

1643 - 1644	SLC19A2:Thiamine-responsive megaloblastic anemia syndrome
	{(NM_006996.3) "c.1223+1G>A", "c.725delC"}
1645 - 1647	SLC1A4:Spastic tetraplegia, thin corpus callosum, and progressive
	microcephaly {(NM_003038.5) "c.1369C>T", "c.766G>A",
1010 1010	"c.944_945del"}
1648 - 1648	SLC22A5:Carnitine deficiency, systemic primary {(NM_003060.3)
	"c.1196G>A"}
1649 - 1650	SLC25A15:Hyperornithinemia-hyperammonemia-homocitrullinemia
	syndrome {(NM_014252) "c.562_564delTTC" (NM_014252.3)
	"c.706A>G"}
1651 - 1652	SLC25A1:Combined D-2- and L-2-hydroxyglutaric aciduria
	{(NM_005984.5) "c.389G>A", "c.845G>A"}
1653 - 1654	SLC25A20:Carnitine-acylcarnitine translocase deficiency - CACT
	{(NM_000387.6) "c.609-3C>G", "c.713A>G"}
1655 - 1655	SLC26A3:Congenital chloride diarhhea (CLD) {(NM_000111.2)
1000	"c.559G>T"}
1656 - 1666	SLC26A4:Pendred syndrome {(NM_000441.2) "c.1001G>T",
	"c.1151A>G", "c.1198delT", "c.1246A>C", "c.1341+1delG",
	"c.1458dupT", "c.2000T>G", "c.2168A>G", "c.349C>T", "c.707T>C",
	"c.716T>A"}
1667 - 1670	SLC29A3:Histiocytosis-lymphadenopathy plus syndrome
	{(NM_018344.5) "c.1157G>A" (NM_018344.6) "c.1045delC",
4674 4670	"c.1279G>A", "c.1309G>A"}
1671 - 1673	SLC2A2:Fanconi-Bickel syndrome {(NM_000340.2) "c.372A>C",
4674 4674	"c.734A>C", "c.901C>T"}
1674 - 1674	SLC30A9:Birk-Landau-Perez cerebro-renal syndrome {(NM_016474.5) "c.1047_1049delCAG"}
1675 - 1676	SLC35A3:Arthrogryposis, mental retardation, and seizures
10/3 - 10/0	{(NM_012243.3) "c.514C>T", "c.886A>G"}
1677 - 1677	SLC35C1:Congenital disorder of glycosylation, type IIc
10// - 10//	{(NM_018389.4) "c.923C>G"}
1678 - 1681	SLC37A4:Glycogen storage disease Ib {(NM_001164277.1)
1070 - 1001	"c.1042_1043delCT", "c.1179G>A", "c.446G>A", "c.83G>A"}
1682 - 1682	SLC39A4:Acrodermatitis enteropathica {(NM_130849.3)
1002 1002	"c.1224delC"}
1683 - 1683	SLC45A2:Albinism, oculocutaneous, type IV {(NM_001012509)
	"c.1076_1077delAG"}
1684 - 1684	SLC46A1:Folate malabsorption, hereditary {(NM_080669)
	"c.337C>T"}
1685 - 1685	SLC4A4:Renal tubular acidosis (RTA), proximal, with ocular
	abnormalities and mental retardation {(NM_003759.3) "c.2321G>A"}
1686 - 1686	SLCO2A1:Hypertrophic osteoarthropathy, primary, Autosomal
	Recessive 2 {(NM_005630.2) "c.1292delC"}
1687 - 1688	SMARCAL1:Schimke immunoosseous dysplasia {(NM_014140.3)
	"c.2542G>T", "c.863-2A>G"}
1689 - 1701	SMPD1:Niemann-Pick disease type B, SMPD1-related {(NM_000543)
	"c.1092-1G>C", "c.880C>A" (NM_000543.5) "c.1493G>T",
	"c.1705T>C", "c.1758T>G", "c.1805G>C", "c.1828_1830delCGC",

	"c.442T>A", "c.573delT", "c.730G>A", "c.911T>C", "c.96G>A",
	"c.996delC"}
1702 - 1702	SNAP29:Cerebral dysgenesis, neuropathy, ichthyosis, and
	palmoplantar keratoderma syndrome - CEDNIK Syndrome
	{(NM_004782) "c.223delG"}
1703 - 1703	SNX10:Osteopetrosis, Autosomal Recessive 8 {(NM_001199835.1)
4704 4707	"c.152G>A"}
1704 - 1707	SPG11:Spastic paraplegia 11, Autosomal Recessive {(NM_025137)
	"c.5986dupT" (NM_025137.4) "c.118C>T", "c.2471dupT", "c.4339C>T"}
1708 - 1712	SPINK5:Netherton syndrome {(NM_001127698.1) "c.2240+5G>A",
1,00 1,12	"c.2557C>T", "c.649C>T", "c.691delC", "c.995delT"}
1713 - 1713	ST3GAL3:Early infantile epileptic encephalopathy 15 {(NM_006279.4)
	"c.958G>C"}
1714 - 1714	STRA6:Microphthalmia {(NM_001142617.1) "c.1678G>C"}
1715 - 1716	STRC:Deafness, Autosomal Recessive 16 {(NM_153700.2)
	"EX7_EX29DEL", "c.4171C>G"}
1717 - 1717	SUCLA2:Mitochondrial DNA depletion syndrome 5 {(NM_003850)
1710 1710	"c.788_802+29del"}
1718 - 1719	SUMF1:Multiple sulfatase deficiency {(NM_182760.3) "c.1043C>T", "c.463T>C"}
1720 - 1721	SURF1:Leigh syndrome, due to COX deficiency {(NM_003172)
1/20 - 1/21	"c.312_321delTCTGCCAGCCinsAT", "c.575_576insTGCG"}
1722 - 1722	SYNE4:Deafness, Autosomal Recessive 76 {(NM_001039876.3)
	"c.228_229delAT"}
1723 - 1723	SZT2:Epileptic encephalopathy, early infantile, 18 {(NM_015284.3)
	"c.73C>T"}
1724 - 1725	SepSecS:Pontocerebellar hypoplasia type 2D {(NM_016955.4)
	"c.1001A>G", "c.715G>A"}
1726 - 1726	TAF2:Mental retardation, Autosomal Recessive 40 {(NM_003184.4) "c.557C>G"}
1727 - 1727	TBCD:Infantile neurodegenerative disorder - Early onset progressive
1,2,1-1,2,	encephalopathy (PEBAT) {(NM 005993.4) "c.1423G>A"}
1728 - 1729	TBCE:Hypoparathyroidism retardation dysmorphism syndrome
	{(NM_003193.5) "c.155_166delGCCACGAAGGGA", "c.355_356del"}
1730 - 1730	TBX19:Adrenocorticotropic hormone deficiency {(NM_005149.3)
	"c.574_577delATAG"}
1731 - 1734	TCIRG1:Osteopetrosis, Autosomal Recessive 1 {(NM_006019.4)
	"c.117+4A>T", "c.1331G>T", "c.1384_1386delAAC", "c.674delG"}
1735 - 1735	TCTN2:Meckel syndrome 8 {(NM_024809.5) "c.1506-2A>G"}
1736 - 1738	TECPR2:Spastic paraplegia 49, Autosomal Recessive {(NM_001172631.2) "c.1319delT", "c.3416delT", "c.566C>T"}
1739 - 1739	TGM1:Ichthyosis, congenital, Autosomal Recessive 1 {(NM_000359)
1/35 - 1/35	"c.2290C>T"}
1740 - 1740	THG1L:Cerebellar ataxia and developmental delay {(NM_017872.5)
	"c.164T>C"}
1741 - 1741	TIMM50:3-methylglutaconic aciduria, type IX {(ENST00000314349.4)
	"c.649C>T"}

1742 - 1744	TK2:Mitochondrial DNA depletion syndrome 2 (myopathic type) {(NM_004614.5) "c.360_361delGCinsAA", "c.361C>A", "c.635T>A"}
1745 - 1745	TKT:Short stature, developmental delay, and congenital heart defects
	{(NM_001135055.2) "c.769_770insCTACCTCCTTATCTTCTG"}
1746 - 1750	TMC1:Deafness, Autosomal Recessive 7 {(NM_138691.2) "c.100C>T",
	"c.1165C>T", "c.1210T>C", "c.1810C>T", "c.1939T>C"}
1751 - 1751	TMEM165:Congenital disorder of glycosylation {(NM_018475.4)
	"c.792+182G>A"}
1752 - 1754	TMEM216:Joubert syndrome 2 (MKS2) {(NM_001173990.3)
	"c.218G>A", "c.218G>T", "c.230G>C"}
1755 - 1755	TMEM231:Meckel syndrome 11 {(NM_001077418.3) "c.664+4A>G"}
1756 - 1756	TMEM260:Neurodevelopmental, Cardiac, and Renal Syndrome
1750 1750	{(NM_017799.3) "c.1393C>T"}
1757 - 1758	TMEM38B:Osteogenesis imperfecta, type XIV {(NM_018112)
1/3/ - 1/30	"c.455_542del", "c.507G>A"}
1750 1761	
1759 - 1761	TMEM67: Joubert syndrome type 6 (MSK3) {(NM_153704)
4-0-4-0-	"c.1065+1delG" (NM_153704.5) "c.1975C>T", "c.725A>G"}
1762 - 1764	TMEM70:ATPase deficiency, nuclear encoded {(NM_017866)
	"c.336T>A" (NM_017866.6) "c.238C>T", "c.316+1G>T"}
1765 - 1766	TMPRSS3:Deafness, Autosomal Recessive 8/10 {(NM_024022)
	"c.1177_1184delins" (NM_024022.2) "c.989delA"}
1767 - 1767	TNNT1:Nemaline myopathy 5, Amish type {(NM_003283)
	"c.574_577delinsTAGTGCTGT"}
1768 - 1768	TPP1:Ceroid lipofuscinosis, neuronal, 2 {(NM_000391) "c.775delC"}
1769 - 1769	TRAK1:Encephalopathy, fatal {(NM_001042646.2) "c.287-2A>C"}
1770 - 1770	TRAPPC9:Mental retardation, Autosomal Recessive 13
2.70	{(NM_031466.7) "c.1423C>T"}
1771 - 1771	TRIM32:Bardet-Biedl syndrome 11 {(NM_012210) "c.388C>T"}
1772 - 1772	TRIOBP:Deafness, Autosomal Recessive 28 {(NM_001039141)
1772-1772	"c.1741C>T"}
1773 - 1774	TRMT10A:Microcephaly, short stature, and impaired glucose
	metabolism {(NM_152292.4) "c.616G>A", "c.727C>T"}
1775 - 1777	TRMU:LIFT, Liver failure infantile transient {(NM_018006)
	"c.500_509del10", "c.835G>A" (NM_018006.5) "c.229T>C"}
1778 - 1781	TRPM1:Night blindness, congenital stationary (complete), 1C,
	Autosomal Recessive {(NM_002420.5) "36.4-KB_DEL,_EX2-7",
	"c.2567G>A", "c.2629C>T", "c.880A>T"}
1782 - 1783	TRPM6:Hypomagnesemia 1, intestinal {(NM_017662.5)
	"c.1010+5G>C", "c.2009+1G>A"}
1784 - 1785	TSHR:Hypothyroidism, congenital, nongoitrous, 1 {(NM_000369)
1707 1703	"c.202C>T" (NM_000369.2) "c.1825C>T"}
1786 - 1786	TSPAN12:Vitroretinal vascular malformations, congenital
1700 - 1700	{(NM_012338) "c.542G>T"}
1707 1707	
1787 - 1787	TSPEAR:Ectodermal dysplasia {(NM_144991.3) "c.1877T>C"}
1788 - 1789	TTN:Cardiomyopathy, dilated - Lethal Congenital Arthrogryposis
1-1-1	{(NM_003319.4) "c.58881dupA" (NM_133432) "c.36122delC"}
1790 - 1790	TUBGCP4:Microcephaly, primary, Autosomal Recessive

\{\text{(NM_014444.5) "c.579dup""} \\ \text{1791 - 1793} \text{TULP1:Retinitis pigmentosa 14 \{\text{(NM_003322) "c.849_852dup" \text{(NM_003322.6) "c.1349G>A", "c.1495+2dup""} \\ \text{1794 - 1795} \text{TYMP:Mitochondrial DNA depletion syndrome 1 \text{(NM_000372) "c.1A>G" \text{(NM_000372.5) "c.1037-1G>A", "c.1037-7T>A", "c.118C>A", "c.1036-A", "c.1037-1G>A", "c.1037-7T>A", "c.118C>A", "c.1204C>T", "c.1217C>T", "c.1357C>T", "c.140G>A", "c.196C>G", "c.454C>T", "c.649C>T", "c.649delC", "c.74dupT", "c.757G>A", "c.332C>T", "c.880G>A", "c.896G>A"} \\ \text{1813 - 1813} \text{UNC13D:Hemophagocytic lymphohistiocytosis, familial, 3 \\ \{\text{(NM_199242.2) "c.753+1G>T"}\} \\ \text{1814 - 1815} \text{UNC80:Hypotonia, infantile, with psychomotor retardation and characteristic facies 2 \(\text{HPR2}\) \{\text{(NM_032504.1) "c.151C>T"}\} \\ \text{1816 - 1816} \text{UPB1:Beta-ureidopropionase deficiency \{\text{(NM_016327.3) "c.917-1G>A"}\} \\ \text{1817 - 1817} \text{UQCRQ:Mitochondrial complex III deficiency, nuclear type 4 \\ \{\text{(NM_014402.5) "c.134C>T"}\} \\ \text{USH16:Usher syndrome, type 1C \{\text{(NM_005709.3) "c.1220delG", "c.238dupC", "c.497-2delA"}\} \\ \text{1821 - 1821} \text{USH16:Usher syndrome, type 1G \{\text{(NM_006933.3) "c.1000C>T", "c.10211delC", "c.12052G>A", "c.1205-2A>G", "c.12575G>A", "c.13274C>T", "c.14020A>G", "c.12067-2A>G", "c.12575G>A", "c.13274C>T", "c.3776elG", "c.3959C>T", "c.4944C>T", "c.509G>A", "c.8519A>C", "c.8519G>T", "c.871G>T", "c.802G>A", "c.8519A>C", "c.9685delG" \\ \text{1848 - 1848} \text{USMG5:Leigh syndrome realted to USMG5 \{\text{(NM_001017535.1) "c.277+1G>T", "c.885C>A"} \\ \text{1149AS39:Arthrogryposis renal dysfunction and cholestasis (ARC) syndrome 2 \{\text{(NM_022067) "c.808C>T"} \\ \text{VPS11:Hypomyelination and developmental delay \{\text{(NM_021729.5) "c.2536T>G"} \\ \text{VPS11:Hypomyelination and developmental delay \{\text{(NM_021729.5) "c.2536T>G"} \\ VPS11:Hypomyelinat
(NM_003322.6) "c.1349G>A", "c.1495+2dupT"} 1794 - 1795
TYMP:Mitochondrial DNA depletion syndrome 1 (MNGIE type) {(NM_001113755.2) "c.433G>A", "c.866A>C"} TYR:Albinism, oculocutaneous, type IA (OCA1A) {(NM_000372)
TYR:Albinism, oculocutaneous, type IA (OCA1A) {(NM_000372)
"c.1A>G" (NM_000372.5) "c.1037-1G>A", "c.1037-7T>A",
"c.1118C>A", "c.1204C>T", "c.1217C>T", "c.1357C>T", "c.140G>A", "c.149C>G", "c.454C>T", "c.649C>T", "c.649delC", "c.74dupT", "c.757G>A", "c.832C>T", "c.880G>A", "c.896G>A"} 1813 - 1813 UNC13D:Hemophagocytic lymphohistiocytosis, familial, 3 {(NM_199242.2) "c.753+1G>T"} 1814 - 1815 UNC80:Hypotonia, infantile, with psychomotor retardation and characteristic facies 2 (HPFR2) {(NM_032504) "c.7183C>T" (NM_032504.1) "c.151C>T"} 1816 - 1816 UPB1:Beta-ureidopropionase deficiency {(NM_016327.3) "c.917-1G>A"} 1817 - 1817 UQCRQ:Mitochondrial complex III deficiency, nuclear type 4 {(NM_014402.5) "c.134C>T"} 1818 - 1820 USH1C:Usher syndrome, type 1C {(NM_005709.3) "c.1220delG", "c.238dupC", "c.497-2delA"} 1821 - 1821 USH1G:Usher syndrome, type 1G {(NM_173477) "c.205dup"} 1822 - 1847 USH2A:Usher syndrome, type 2A {(NM_206933.3) "c.1000C>T", "c.10211delC", "c.12052G>A", "c.12067-2A>G", "c.124575G>A", "c.13274C>T", "c.14020A>G", "c.14023A>T", "c.14413G>A", "c.519G>T", "c.5776+1G>A", "c.236_239dupGTAC", "c.3368A>G", "c.377delG", "c.3959C>T", "c.464C>T", "c.5078G>A", "c.8558+1G>T", "c.8719A>C", "c.9685delG"} 1848 - 1848 USMG5:Leigh syndrome realted to USMG5 {(NM_032747) "c.87+1G>C"} "c.87+1G>C"} 1849 - 1850 VDR:Rickets, vitamin D-resistant, type IIA {(NM_001017535.1) "c.277+1G>T", "c.885C>A"} VIPAS39:Arthrogryposis renal dysfunction and cholestasis (ARC) syndrome 2 {(NM_022067) "c.808C>T"}
"c.149C>G", "c.454C>T", "c.649C>T", "c.649delC", "c.74dupT", "c.757G>A", "c.832C>T", "c.880G>A", "c.896G>A"} 1813 - 1813
"c.757G>A", "c.832C>T", "c.880G>A", "c.896G>A"} 1813 - 1813
1813 - 1813
\text{\{NM_199242.2\}\] "c.753+1G>T"\} \
UNC80:Hypotonia, infantile, with psychomotor retardation and characteristic facies 2 (HPFR2) {(NM_032504) "c.7183C>T" (NM_032504.1) "c.151C>T"}
characteristic facies 2 (HPFR2) {(NM_032504) "c.7183C>T"
(NM_032504.1) "c.151C>T"} 1816 - 1816
1816 - 1816 UPB1:Beta-ureidopropionase deficiency {(NM_016327.3) "c.917-1G>A"} 1817 - 1817 UQCRQ:Mitochondrial complex III deficiency, nuclear type 4 {(NM_014402.5) "c.134C>T"} 1818 - 1820 USH1C:Usher syndrome, type 1C {(NM_005709.3) "c.1220delG", "c.238dupC", "c.497-2delA"} 1821 - 1821 USH1G:Usher syndrome, type 1G {(NM_173477) "c.205dup"} 1822 - 1847 USH2A:Usher syndrome, type 2A {(NM_206933.3) "c.1000C>T", "c.10211delC", "c.12052G>A", "c.12067-2A>G", "c.12575G>A", "c.13274C>T", "c.14020A>G", "c.14023A>T", "c.14413G>A", "c.13274C>T", "c.14020A>G", "c.14023A>T", "c.14413G>A", "c.14424C>A", "c.2167+5G>A", "c.2209C>T", "c.4544C>T", "c.5078G>A", "c.5519G>T", "c.5776+1G>A", "c.6159delA", "c.6937G>T", "c.802G>A", "c.8558+1G>T", "c.8719A>C", "c.9685delG"} 1848 - 1848 USMG5:Leigh syndrome realted to USMG5 {(NM_032747) "c.87+1G>C"} 1849 - 1850 VDR:Rickets, vitamin D-resistant, type IIA {(NM_001017535.1) "c.277+1G>T", "c.885C>A"} 1851 - 1851 VIPAS39:Arthrogryposis renal dysfunction and cholestasis (ARC) syndrome 2 {(NM_022067) "c.808C>T"} 1852 - 1852
1G>A"} 1817 - 1817 1818 - 1820 1818 - 1820 USH1C:Usher syndrome, type 1C {(NM_005709.3) "c.1220delG", "c.238dupC", "c.497-2delA"} 1821 - 1821 USH1G:Usher syndrome, type 1G {(NM_173477) "c.205dup"} 1822 - 1847 USH2A:Usher syndrome, type 2A {(NM_206933.3) "c.1000C>T", "c.10211delC", "c.12052G>A", "c.12067-2A>G", "c.12575G>A", "c.13274C>T", "c.14020A>G", "c.14023A>T", "c.14413G>A", "c.14424C>A", "c.2167+5G>A", "c.2209C>T", "c.236_239dupGTAC", "c.3368A>G", "c.377delG", "c.3959C>T", "c.4544C>T", "c.5078G>A", "c.5519G>T", "c.5776+1G>A", "c.6159delA", "c.6937G>T", "c.802G>A", "c.8558+1G>T", "c.8719A>C", "c.9685delG"} 1848 - 1848 USMG5:Leigh syndrome realted to USMG5 {(NM_032747) "c.87+1G>C"} 1849 - 1850 VDR:Rickets, vitamin D-resistant, type IIA {(NM_001017535.1) "c.277+1G>T", "c.885C>A"} VIPAS39:Arthrogryposis renal dysfunction and cholestasis (ARC) syndrome 2 {(NM_022067) "c.808C>T"} 1852 - 1852 VPS11:Hypomyelination and developmental delay {(NM_021729.5)
1817 - 1817
Section Sect
"c.238dupC", "c.497-2delA"} 1821 - 1821 USH1G:Usher syndrome, type 1G {(NM_173477) "c.205dup"} USH2A:Usher syndrome, type 2A {(NM_206933.3) "c.1000C>T", "c.10211delC", "c.12052G>A", "c.12067-2A>G", "c.12575G>A", "c.13274C>T", "c.14020A>G", "c.14023A>T", "c.14413G>A", "c.14424C>A", "c.2167+5G>A", "c.2209C>T", "c.236_239dupGTAC", "c.3368A>G", "c.377delG", "c.3959C>T", "c.4544C>T", "c.5078G>A", "c.5519G>T", "c.5776+1G>A", "c.6159delA", "c.6937G>T", "c.802G>A", "c.8558+1G>T", "c.8719A>C", "c.9685delG"} 1848 - 1848 USMG5:Leigh syndrome realted to USMG5 {(NM_032747) "c.87+1G>C"} VDR:Rickets, vitamin D-resistant, type IIA {(NM_001017535.1) "c.277+1G>T", "c.885C>A"} 1851 - 1851 VIPAS39:Arthrogryposis renal dysfunction and cholestasis (ARC) syndrome 2 {(NM_022067) "c.808C>T"} 1852 - 1852 VPS11:Hypomyelination and developmental delay {(NM_021729.5)
1821 - 1821 USH1G:Usher syndrome, type 1G {(NM_173477) "c.205dup"} 1822 - 1847 USH2A:Usher syndrome, type 2A {(NM_206933.3) "c.1000C>T", "c.10211delC", "c.12052G>A", "c.12067-2A>G", "c.12575G>A", "c.13274C>T", "c.14020A>G", "c.14023A>T", "c.14413G>A", "c.14424C>A", "c.2167+5G>A", "c.2209C>T", "c.236_239dupGTAC", "c.3368A>G", "c.377delG", "c.3959C>T", "c.4544C>T", "c.5078G>A", "c.5519G>T", "c.5776+1G>A", "c.6159delA", "c.6937G>T", "c.802G>A", "c.8558+1G>T", "c.8719A>C", "c.9685delG"} 1848 - 1848 USMG5:Leigh syndrome realted to USMG5 {(NM_032747) "c.87+1G>C"} VDR:Rickets, vitamin D-resistant, type IIA {(NM_001017535.1) "c.277+1G>T", "c.885C>A"} 1851 - 1851 VIPAS39:Arthrogryposis renal dysfunction and cholestasis (ARC) syndrome 2 {(NM_022067) "c.808C>T"} 1852 - 1852 VPS11:Hypomyelination and developmental delay {(NM_021729.5)
1822 - 1847 USH2A:Usher syndrome, type 2A {(NM_206933.3) "c.1000C>T", "c.10211delC", "c.12052G>A", "c.12067-2A>G", "c.12575G>A", "c.13274C>T", "c.14020A>G", "c.14023A>T", "c.14413G>A", "c.14424C>A", "c.2167+5G>A", "c.2209C>T", "c.236_239dupGTAC", "c.3368A>G", "c.377delG", "c.3959C>T", "c.4544C>T", "c.5078G>A", "c.5519G>T", "c.5776+1G>A", "c.6159delA", "c.6937G>T", "c.802G>A", "c.8558+1G>T", "c.8719A>C", "c.9685delG"} 1848 - 1848 USMG5:Leigh syndrome realted to USMG5 {(NM_032747) "c.87+1G>C"} 1849 - 1850 VDR:Rickets, vitamin D-resistant, type IIA {(NM_001017535.1) "c.277+1G>T", "c.885C>A"} 1851 - 1851 VIPAS39:Arthrogryposis renal dysfunction and cholestasis (ARC) syndrome 2 {(NM_022067) "c.808C>T"} 1852 - 1852 VPS11:Hypomyelination and developmental delay {(NM_021729.5)
"c.10211delC", "c.12052G>A", "c.12067-2A>G", "c.12575G>A",
"c.13274C>T", "c.14020A>G", "c.14023A>T", "c.14413G>A",
"c.14424C>A", "c.2167+5G>A", "c.2209C>T", "c.236_239dupGTAC",
"c.3368A>G", "c.377delG", "c.3959C>T", "c.4544C>T", "c.5078G>A",
"c.5519G>T", "c.5776+1G>A", "c.6159delA", "c.6937G>T", "c.802G>A", "c.8558+1G>T", "c.8719A>C", "c.9685delG"} 1848 - 1848
"c.8558+1G>T", "c.8719A>C", "c.9685delG"} 1848 - 1848 USMG5:Leigh syndrome realted to USMG5 {(NM_032747)
1848 - 1848 USMG5:Leigh syndrome realted to USMG5 {(NM_032747)
"c.87+1G>C"} 1849 - 1850 VDR:Rickets, vitamin D-resistant, type IIA {(NM_001017535.1)
1849 - 1850 VDR:Rickets, vitamin D-resistant, type IIA {(NM_001017535.1)
"c.277+1G>T", "c.885C>A"} 1851 - 1851 VIPAS39:Arthrogryposis renal dysfunction and cholestasis (ARC) syndrome 2 {(NM_022067) "c.808C>T"} 1852 - 1852 VPS11:Hypomyelination and developmental delay {(NM_021729.5)
syndrome 2 {(NM_022067) "c.808C>T"} 1852 - 1852 VPS11:Hypomyelination and developmental delay {(NM_021729.5)
1852 - 1852 VPS11:Hypomyelination and developmental delay {(NM_021729.5)
"c 2536T>C"\
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1853 - 1856 VPS13A:Choreoacanthocytosis {(NM_033305.3) "c.2343delA",
"c.6059delC", "c.9446_9449dup", "delexon70-73"}
1857 - 1858 VPS13B:Cohen syndrome {(NM_017890) "c.4894C>T"
(NM_017890.4) "c.6732+1G>A"}
VPS33B:Arthrogryposis renal dysfunction and cholestasis (ARC) syndrome {(NM_018668.4) "c.403+1G>A", "c.700G>C"}
1861 - 1861 VPS37A:Spastic paraplegia 53, Autosomal Recessive {(NM_152415.3)
"c.1146A>T"}
1862 - 1862 VPS45:Neutropenia, severe congenital, 5, Autosomal Recessive
{(NM_007259.5) "c.671C>A"}

	{(NM_001128159.3) "c.1556+5G>A", "c.2084A>G"}
1865 - 1865	VRK1:Pontocerebellar hypoplasia type 1A {(NM_003384.3) "c.1072C>T"}
1866 - 1868	WFS1:Wolfram-like syndrome, Autosomal Dominant {(NM_006005) "c.1230_1233delCTCT", "c.1770_1773delGTCT", "c.2590G>A"}
1869 - 1870	WISP3:Arthropathy, progressive pseudorheumatoid, of childhood {(NM_003880.3) "c.156C>A ", "c.536_537delGT"}
1871 - 1871	XPC:Xeroderma pigmentosum, group C {(NM_004628.4) "c.566_567delAT"}
1872 - 1872	XRCC2:Fanconi Anemia {(NM_005431.1) "c.643C>T"}
1873 - 1873	ZBTB24:Immunodeficiency-centromeric instability-facial anomalies syndrome-2 {(NM_014797.2) "c.501dupA"}
1874 - 1874	ZMPSTE24:Mandibuloacral dysplasia with type B lipodystrophy {(NM_005857) "c.1085dupT"}
1875 - 1876	ZNF469:Brittle cornea syndrome 1 {(NM_001127464.2) "c.5943delA", "c.9531delG"}

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לפי החלטת איגוד הגנטיקאים הישראלי ,המוטציה מדווחת חיובית רק ליהודים ממוצא קווקזי, De boer M, Gavrieli R, Van leeuwen K, et al. A false-carrier state for the c.579G>A mutation in the NCF1 gene in Ashkenazi Jews. J Med Genet. 2018;55(3):166-172.

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לפי החלטת איגוד הגנטיקאים הישראלי ,המוטציה מדווחת חיובית רק לאנשים ממוצא דרוזי, Belostotsky R, Seboun E, Idelson GH, et al. Mutations in DHDPSL are responsible for primary hyperoxaluria type III. Am J Hum Genet. 2010;87(3):392-399. doi:10.1016/j.ajhg.2010.07.023.